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Specially Revised and Enlarged by the Author from the
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Vorrede zur 4. englischen Ausgabe.

Der plötzliche Tod meines verehrten Freundes Dr. James Cagney zwang mich, nachdem die 3. englische Auflage in der kürzesten Zeit vergriffen war, die Revision und Ergänzung der 4. englischen Auflage *selbst* zu besorgen.

Es erfüllt mich mit grossem Stolze, dass das Buch gerade bei den englischen Collegen eine so grosse Verbreitung gewonnen hat und hoffe ich, dass es auch in dieser Ausgabe die Gunst der alten Freunde sich erhalten und neue hinzu erwerben wird.

Leider ist es mir versagt, meinem verewigten Freunde *Cagney* an dieser Stelle zu danken, um so mehr gedenke ich dankbar der Dienste, welche der Verewigte diesem Buche geleistet hat!

PREFACE TO THE FOURTH ENGLISH EDITION.

Owing to the sudden death of my esteemed friend Dr. James Cagney, I have been obliged, in consequence of the rapidity with which the third English edition of this work was exhausted, to undertake the work of revising and supplementing the fourth English edition myself.

It is a source of pride to me to find that the work has met with such an extensive circulation among my English colleagues, and I hope that this new edition will both meet with favour from old friends and also lead to the acquisition of new ones.

Unhappily, all opportunity of thanking my deceased friend Cagney has now passed away; so much the more then do I hold in grateful remembrance the services rendered by him in connection with this work.

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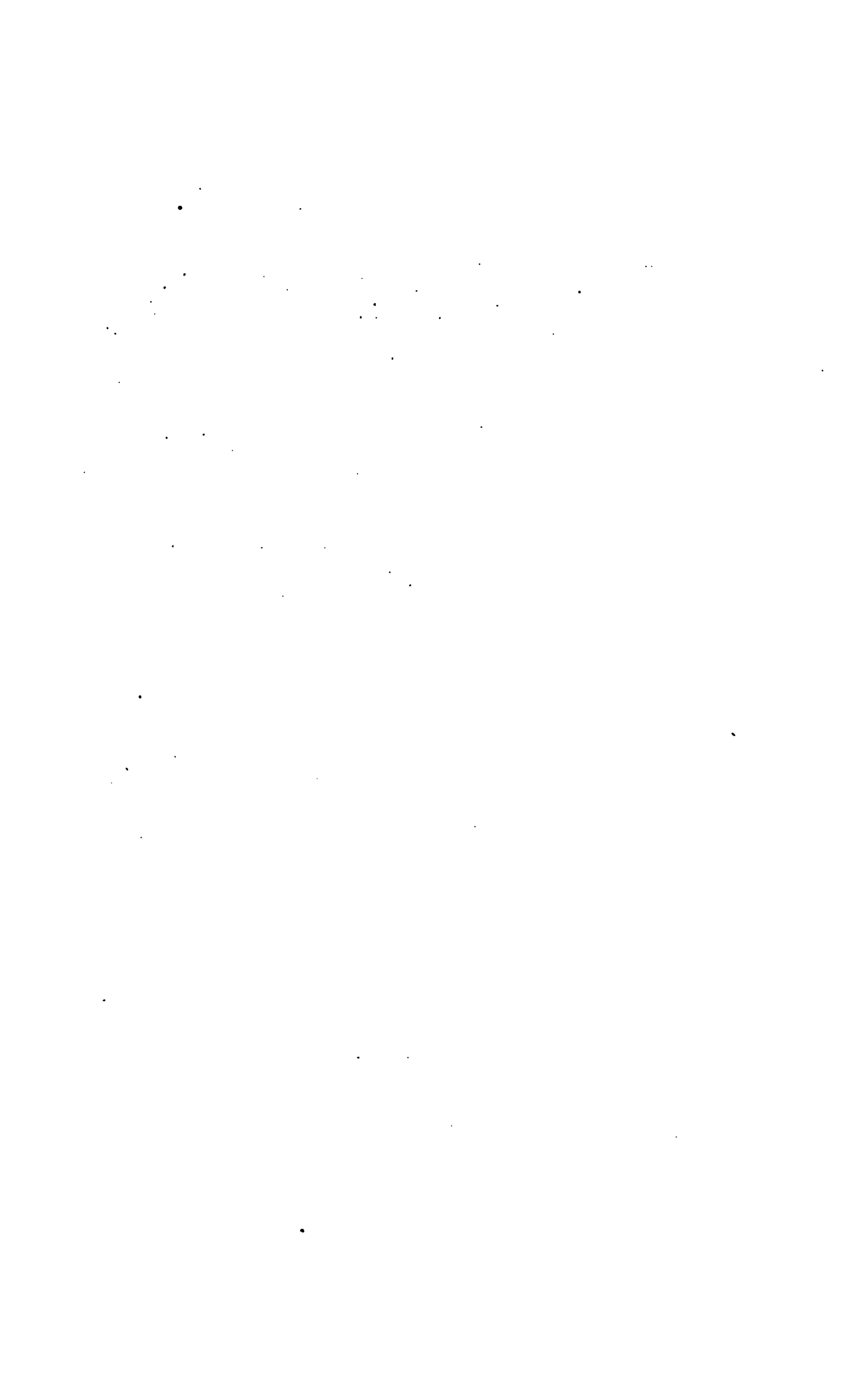


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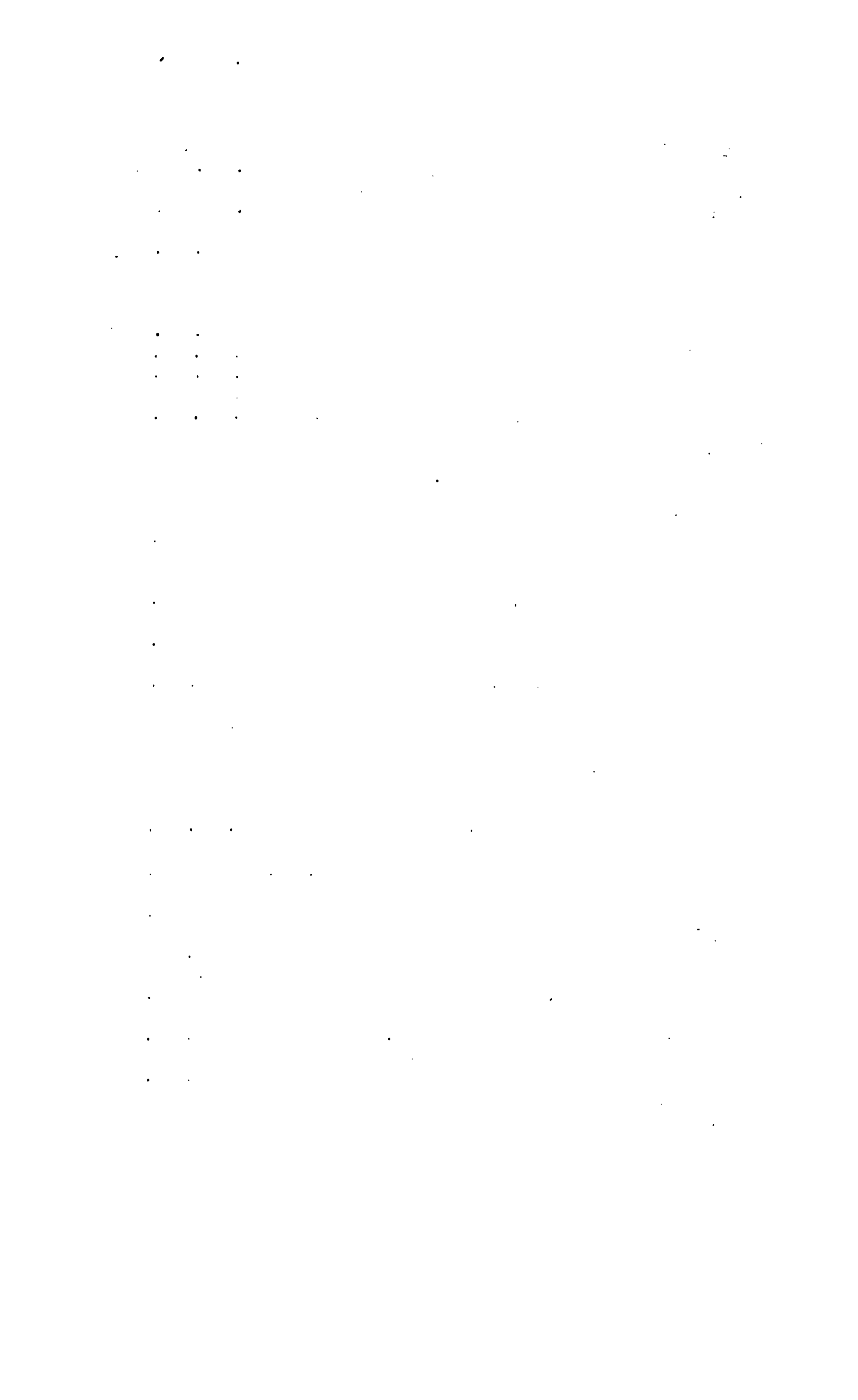
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CLINICAL DIAGNOSIS

CHAPTER I

THE BLOOD

EVERY change in the quantity or the quality of the blood itself is apt to be attended with serious disturbance of the system ; further, the blood is the carrier and distributer through the body of nearly all the poisons, organic and inorganic, which act upon the latter. From this it follows that the physiology and pathology of the blood represent a mass of knowledge at once immense and various. It is not our purpose here, however, to treat these subjects exhaustively, but merely to select from them certain well-established facts which bear upon disease, and to point out the way in which they may help in its diagnosis.

I. COLOUR.—Arterial and venous blood differ considerably as to colour in health, the former being scarlet, and the latter a bluish red. The distinction, however, belongs not to the fluid part of the blood, or plasma, but to the colouring-matter, or hæmoglobin, contained in the red corpuscles, and it depends upon the chemical constitution of the corpuscles, which, changing colour themselves, determine the tint of the whole mass of the blood.

Thus, when the blood is rich in oxygen, the amount of hæmoglobin is increased, and the fluid is proportionately bright red. Again, where, as is always the case with venous blood, oxygen is deficient, or where, from physiological or pathological causes, arterial blood contains but little oxyhæmoglobin, the colour is darker, and this in a degree corresponding to the condition which underlies it. In certain morbid states, however, the blood may assume a brighter tint than that of healthy arterial blood. In carbonic oxide poisoning, for instance, it is of a bright cherry colour (see p. 76). [Venous blood, which appears dark red by reflected light, is green by transmitted light. It is therefore said to be dichroic. Arterial blood is monochroic.^{1]}* The blood which

* The Numbers refer to the Bibliography at the end of the volume.

is taken from the finger for the purpose of microscopical examination by means of a slight puncture is usually venous in character.

II. THE REACTION of healthy blood, like that of all the tissue fluids, except the urine and gastric juice, is alkaline. [It is due to the presence of disodic phosphate, Na_2HPO_4 , and bicarbonate of soda.] Still, this reaction is liable to considerable variation under certain conditions of health as well as in disease.

The alkalinity of the blood diminishes when it has been withdrawn from the influence of living blood-vessels. Hence we find that an acid reaction is one of the phenomena of coagulation, and that this becomes more pronounced the longer the blood is allowed to stand. [The change depends upon the formation of an acid derived, probably, from the decomposition of the colouring-matter of the red corpuscles.]

To test the reaction of the blood, *Liebreich*² employs plates of plaster of Paris or clay soaked in neutral litmus solution. A few drops of the blood to be examined are placed upon this, and washed off again with water.* If the blood was alkaline, the spot upon which it has fallen exhibits a blue, but if acid, a red colour.

For the same purpose *Zuntz*³ uses glazed litmus paper soaked in a solution of common salt or sulphate of soda ; this he dips several times in the blood to be tested, and again washes in the salt solution. The same thing may be done by allowing a drop of the blood to fall on the litmus paper, and then quickly washing it off again, as in Liebreich's method.

For the comparative estimation of the alkalinity of the blood in animals, *Lassar*⁴ has devised a process, which, however, cannot be applied to the case of human beings, where the requisite quantity of blood is not available. On the other hand, the plan which *Landois*⁵ recommends is very suitable for clinical purposes.

The author has obtained useful results in a large series of observations by proceeding according to the following method (a modification of that of Landois), for the quantitative estimation of the alkalinity of the blood :—

A mixture is made of a concentrated solution of sulphate of soda with $1/100$ and $1/1000$ normal solution of tartaric acid (I.) in various proportions ; and in this way a series of test-fluids is obtained, the members of which contain varying quantities of acid to the cubic centimetre.†

* [The reaction of the blood with ordinary litmus paper is obscured by the red colour of the fluid, and the various expedients for its determination are directed to overcome this difficulty.]

† The fluids are prepared thus : 7.5 grms. of pure tartaric acid are dissolved in a litre of water, and a normal solution ($1/10$ of the acid) results. By appropriate dilution of this, the other normal solutions ($1/100$ and $1/1000$) may be obtained.⁶

REACTION OF THE BLOOD

3

Experience shows that eighteen such test-solutions of varying acidity are needed. And of these :—

I.	contains in 1 cc. 0.9 cc. of 1/100 normal solution of acid and 0.1 cc.						
II.	,, in 1 cc. 0.8 cc. of 1/100	„	„	„	0.2 „		
	and so on.						
IX.	„ in 1 cc. 0.1 cc. of 1/100	„	„	„	0.9 „		
X.	„ in 1 cc. 0.9 cc. of 1/1000	„	„	„	0.1 „		
	and so on.						
XIV.	„ in 1 cc. 0.5 cc. of 1/1000	„	„	„	0.5 „		
	and so on.						
XVIII.	„ in 1 cc. 0.1 cc. of 1/1000	„	„	„	0.9 „		

The experiment is conducted in the following manner :—The proper quantities of the acid and sulphate of soda solutions are placed in a series of watch-glasses, by means of a pipette graduated in 0.1 cc.,* and a number of strips of very sensitive blue and red litmus paper are prepared. To make these, filter paper is saturated with the litmus solution prepared according to *Mays'*⁷ prescription, dried, cut into strips, and applied according to the object in view. The blood is usually taken by means of cupping-glasses from the patient's back, and, before it coagulates, 0.1 cc. of the blood is added to each cc. of the fluids described above, well mixed in each case, and the resulting mixtures tested with the litmus papers until one is found to exhibit a neutral reaction, *i.e.* leaving the red and blue litmus paper unchanged. This will show what quantity of the acid is required to neutralise 0.1 cc. of the blood in question. In order to arrive at an accurate result, it is necessary to proceed very quickly, and as a rule it may be laid down that not more than 1½ minutes should be allowed to elapse between the taking of the blood and the conclusion of the experiment, having regard to the rapid diminution of alkalinity after the blood is withdrawn from the living vessels.

For the sake of clearness the following example may be taken :—

In the case of a man who suffered from tuberculosis and tabes dorsalis, it was found that 0.4 cc. of 1/100 normal tartaric acid solution was required to neutralise 0.1 cc. of blood.

1 cc. of 1/100 normal solution of the acid corresponds to 0.0004 grm. NaOH.

O. I	"	"	"	"	"	0.00004	"	"
O. 4	"	"	"	"	"	0.00016	"	"

The alkalinity of the blood may therefore be expressed by 0.00016 grm. NaOH for 0.1 cc., or .160 grm. NaOH for 100 cc.

Haycraft and Williamson's Method.

[The method recently introduced by *Haycraft* and *Williamson*⁸ is very suitable for clinical purposes, since by its means the alkalinity of the blood may be determined quantitatively from a single drop of the fluid.

* For this purpose also the automatic pipettes (see p. 23) containing 0.1 cc. will serve very well.

A number of red litmus papers is prepared, containing varying quantities of oxalic or some other acid. One of these is such as is found by experiment barely to give a reaction with normal blood, and this is made the basis of a series of test papers answering to different degrees of alkalinity. The strength of each is estimated by means of a solution of caustic potash of known concentration. The papers are glazed, and dipped for a second or two in liquid paraffin and then dried. In conducting the experiment a drop of blood is drawn from the finger (previously well cleansed) and placed upon a paper of medium strength. There it is allowed to rest for ten seconds. Sufficient of the plasma has then soaked in. The blood is washed off, and the reaction, if any, is at once apparent. Should this be so, a paper containing more acid is employed; but if there be no reaction, a weaker one is taken. Suppose, now, it is found that the blood will give a reaction with the sixth and not with the seventh paper, the former is then taken as the expression of its alkalinity. But it is known that an $\frac{n}{x}$ solution of an alkali

will give the same reaction, therefore the alkalinity of the blood will be $\frac{n}{x}$. "This is perhaps not absolutely true, for probably the blood plasma does not percolate so readily into the litmus paper as does a watery solution of an alkali. In this case, however, the error will be uniform," and will not vitiate the conclusion in a series of comparative investigations. The objection thus anticipated is insisted on by *Hutchinson*,⁸ who points out that by *Haycraft's* method the alkalinity of the blood in anaemia is apparently increased. For the explanation of this, see p. 94. The statement in the text concerning the observations of *A. Loewy* and of *Schultz-Schultzenstein* would seem to bear upon this point. *Hutchinson* prefers the titration method, on the ground that the other, the percolation test, takes insufficient account of the alkali within the corpuscles.]

*Tauszki*⁹ weighs the blood used, and titrates with tropæolin or litmus. He professes to get good results in this way.

It is true, as *H. Meyer*¹⁰ has shown, that the results to be obtained in this way are open to error. This, indeed, we should expect, since the final reaction varies in each specimen with the colour of the blood and the quantity of CO₂ which it contains. The method has been described here, faulty and unsatisfactory as it is, because by means of it certain information has been obtained as to the character of the blood in disease.¹¹

The observations of *A. Loewy*¹² in the titration of blood with litmus, and of *Schultz-Schultzenstein*,¹³ who employed the titration method with erythrosin as indicator and the blood dissolved in water, tend to show that the alkalinity of the blood is really greater than the method given would show it would be; and although it is possible that the results obtained may require correction later, the author's experiments¹⁴ have led him to the conclusion that the alkalinity of 100 cc. of healthy human blood corresponds to 260–300 mgrms. of NaOH. *Canard*,¹⁵ who adopts a similar method, gives the equivalent as 203–276 mgrms. NaOH; while *Mya* and *Tassinari*,¹⁶ from experiments upon blood drawn from the veins, quote very much higher figures (516 mgrms.). The alkalinity of the blood is often diminished in fever. The author

has invariably found it reduced considerably in uræmia, as well as in certain toxic states, as in carbonic oxide, and especially in phosphorus poisoning.¹⁷ [It is also reduced in persistent vomiting.] In organic disease of the liver, leukæmia, pernicious anaemia, and diabetes, the author has found such a diminution as was capable of being expressed in figures, and his views (the condition of the blood in chlorosis alone excepted) are borne out by the researches of *Grüber*.¹⁸ His conclusions have also been confirmed by *Peiper*¹⁹ and *Rumpf*.²⁰ The results obtained by *Kraus*²¹ with another method are substantially the same.

The blood is taken by a lancet, and its contents of carbonic acid determined by weighing. This method is probably accurate; but from the quantity of blood which it demands, and the inexpediency of employing the lancet for clinical purposes, it is not to be preferred to the other methods, which in general have yielded the same results.

*Klemperer*²² has proceeded also on the principle of estimating the proportion of CO₂. He has found that the diminished alkalinity of the blood in fever is not affected by the administration of anti-pyretic remedies. *Cantani*²³ is of opinion that the blood in cholera may exhibit an acid reaction even during life.

III. SPECIFIC GRAVITY OF THE BLOOD.—The specific gravity of healthy human blood has been stated at 1.045–1.075 by *Landois*,²⁴ and 1.035–1.068 by *Lloyd Jones*.²⁵ It is usually lower in women than in men; and the last-mentioned authority has shown that it is very high at birth, reaching to 1.056–1.066.²⁶ It then falls gradually during the first few years of life, and reaches its maximum in man at 35–45 years of age. [It is diminished by hunger, in pregnancy, by the ingestion of solid or liquid food, or by gentle exercise.²⁷]

To estimate the specific gravity of the blood, *Roy's method*, as used by *Devoto* and *Siegl*, may be adopted. This requires a series of test-tubes, holding a mixture of glycerine and water in different proportions, so that the sp. gr. of these shall range between 1.040 and 1.080. The test-tubes should have a diameter of 4 cm. and should hold from 80–100 cc.

The proceeding is as follows:²⁸—A drop of blood is drawn from the finger by pricking it with an aseptic needle; a capillary tube of glass bent at a right angle, and connected by a caoutchouc tube with the nozzle of a Pravaz syringe, is now taken, and the free pointed end of the tube is placed in the centre of the exuding blood, some of which is drawn into the tube by a slight movement of the piston. By gentle pressure on the piston of the syringe a drop of blood is expelled into the middle of the fluid in one of the test-tubes. There the blood will rise or sink according to the density of the glycerine mixture, and successive trials

are made until a fluid is found in which the blood remains suspended. The density of this fluid is that of the blood. The glycerine mixtures may be preserved for future use by the addition of a little thymol, but their sp. gr. must be verified before each investigation. [English observers generally dispense with the syringe and employ the right-angled glass-tube, which is pointed at one extremity and expanded at the other, where it is closed by a caoutchouc cap, and by means of this the blood is drawn in and again expressed in the test fluid. For purposes of comparison the blood should be examined in the morning, and always at the same hour, since *Lloyd Jones*²⁹ has shown that its sp. gr. undergoes diurnal variations, dependent probably upon the ingestion of food. Again, in removing the blood for examination, it is important to avoid pressure on the part from which the blood is withdrawn (as by squeezing, or the application of a ligature), since the sp. gr. is altered by such expedients.]

[Instead of glycerine *Landois* employs solutions of sulphate of soda for the test fluids. The process is in other respects the same as *Roy's*.³⁰

Monckton Copeman and *Sherrington*³¹ have investigated the sp. gr. of the blood by a method founded on that of Roy. To prevent decomposition of the test fluids and consequent change in their density they are derived from a stock fluid of glycerine and water saturated with boro-glyceride and sulphate of magnesium, with a small quantity—1 in 1000—of corrosive sublimate. Such a fluid will remain serviceable for more than three years.

Hammerschlag's method for determining the specific gravity of the blood is a modification of Roy's, with the advantage that it can be much more rapidly and conveniently applied. A mixture is made, in a suitable vessel, of chloroform and benzol, two liquids of widely differing specific gravity and which are freely miscible. In this mixture a drop of blood is placed, and by the addition as required of more chloroform or benzol, the density of the liquid is altered until the blood remains suspended in it. When this point is reached, the specific gravity of the liquid is ascertained in the usual way. That of the blood is of course the same. Chloroform and benzol may be supplied from two pipettes, and the whole proceeding may be accomplished in a few seconds. This method has given the most satisfactory results; and it dispenses with the troublesome necessity of preparing a series of solutions.^{32]}

It is the author's opinion that *Hammerschlag's*³³ method, and that of *Schmaltz*³⁴ and *Peiper*,³⁵ who make use of a capillary pycnometer, seem to have no advantage over that of Roy.

Examination of the blood in the author's clinic has shown that intestinal haemorrhage, severe anaemia, and prostration are attended with a fall in specific gravity. *Siegl* and *Schmaltz* have found that the specific gravity varies with the proportion of haemoglobin, but not with the total

number of cellular elements. Hence it follows that a diminution in the amount of haemoglobin may be inferred from a fall in sp. gr., and the comparatively easy investigation of the latter may be made in practice to replace the estimation of haemoglobin, which can be effected only by the use of costly instruments. According to *Monti*,³⁶ the density of the blood in childhood shows some peculiarities.

IV. CHANGES IN THE FORMED ELEMENTS OF THE BLOOD.

—The blood contains red and white corpuscles, and recent observations (*Bizzozero*) have shown the presence in it of a third class of formed elements,—the (blood-tablets or blood-plates) (fig. 1). The existence of these bodies is now beyond dispute. To make them apparent in fresh blood, it is necessary to fix the latter by the addition of some preserving fluid, such as *Hayem's*³⁷ solution, when it may be examined directly with an oil-immersion lens and a narrow diaphragm.

The constitution of *Hayem's solution* is as follows :—

1 grm. of chloride of sodium, .5 grms. of sodic sulphate, 0.5 grm. corrosive sublimate, and 200 grms. distilled water.

The preparation will then show the bodies in question as minute objects with a diameter less than half that of the red blood-corpuscles, scattered singly, or in groups in the field. In the present state of our knowledge, they possess no diagnostic importance. [They are supposed to be most abundantly present in the blood of persons suffering from chronic diseases ; and are said to increase in number during pregnancy (*Halla*), in conditions of regeneration (*Afanassiew*), in febrile anaemia (*Fusari*), and to diminish in fever.³⁸]

To estimate their number the same instrument may be used as for the enumeration of leucocytes (p. 13). To dilute the blood *Pruss*³⁹ uses a modified Fleming's fluid, a mixture of chromic, acetic, and osmic acids. The tendency of the blood-plates to adhere may be met by the addition of peptone in a solution of common salt coloured with methyl violet.

*H. Rabl*⁴⁰ recommends the employment of the iron-haematoxylin method of staining proposed by Heidenhain for revealing the centrosomes. The air-dry preparations are fixed in a $\frac{3}{4}$ per cent. solution of common salt, saturated with sublimate solution, the cover-glass preparations being allowed to remain in the liquid for a quarter to half an hour. They are then thoroughly washed with water and immersed for an hour in a one-half per cent. solution of iron alum, again washed with distilled water, and left for half an hour in a freshly prepared aqueous solution of haematoxylin. When all the elements in the preparation have become blue-black in colour it is immersed in a very dilute solution of iron alum until the preparation has assumed a greenish-yellow coloration, whereupon it is washed with distilled water, dried, and mounted. In

these preparations the blood-plates and leucocytes are stained blue-black, whilst the red blood-corpuses are decolorised. It is highly desirable to subsequently stain the latter with picric acid or aurantia.

The physiology of the red and white blood-corpuses is sufficiently set forth in the Text-books of that science.⁴¹

Pathologically, the corpuses exhibit changes as to quantity and character which are of the utmost importance in diagnosis. These changes seldom occur separately, but are usually combined—although alterations of the *structure* of the corpuses may be more pronounced in some cases, of their *number* in others. We shall consider—

1. The *diminution* in the number of the cellular elements of the blood (*oligocythaemia*)
2. The *increase* in the number of the cellular constituents. An



FIG. 1. Blood-plates from normal blood. (The blood had been fixed with Hayem's solution. Eye-piece 16*L*, objective Zeiss $\frac{1}{2}$, homogeneous immersion.)

absolute increase of the red corpuses (*polycythaemia rubra transitoria*) has been shown by *Taussig*⁴² and *v. Jakob*⁴³ to occur in phosphorus poisoning; by *Wolf*⁴⁴ and others as a normal event at the menopause. *J. K. Mitchell*⁴⁵ has found that massage increases the number of red corpuses, and also the quantity of haemoglobin. A relative preponderance of white corpuses is often met with. This happens normally during digestion (*physiological leucocytosis*)—as a transient phenomenon in a number of morbid states (*temporary leucocytosis*), and as a persistent condition (*leukemia* and *pathological leucocytosis*).

3. Changes in the form of the blood-corpuses (*poikilocytosis, microcythaemia*).

4. Changes in the size of the corpuses, and especially of the red corpuses.

1. Oligocythæmia.—*Vierordt* has computed that in health the number of red blood-corpuscles is five millions in a man, and $4\frac{1}{2}$ millions in a woman to the cubic millimetre of blood.^{46,47} In disease the number may diminish temporarily or permanently to two millions, or even sink as low as 360,000 per cubic millimetre. Such a condition may occur as a consequence of haemorrhage, whether of a traumatic origin, or due to morbid changes in the blood-vessels, as when intestinal bleeding takes place in typhoid fever, or bleeding from the stomach in gastric ulcer, or from the oesophagus. As a permanent state, it may be a phenomenon of any disease which is attended with deficient regeneration of the blood.

[The number of the red corpuscles is lessened in chronic lead-poisoning, miasmatic conditions, and in syphilis.]

Diagnosis of Oligocythæmia.—The methods and apparatus employed by physiologists to estimate a diminution of the red blood-corpuscles are very many ; but of these a large number are useless for clinical purposes, inasmuch as they require too great quantities of blood to work upon.

The apparatus which will serve our purpose are of two classes. One is used to count the actual number of blood-corpuscles in a specimen of blood ; the other, by estimating the quantity of haemoglobin present, enables us to draw an inference as to changes of the blood. Both methods have their advantages and supplement each other, since a diminution in the haemoglobin is usually proportionate to a diminution in the red corpuscles ; thus, *oligochromæmia* and *oligocythæmia* mostly occur together. Quite recently also, it has been found possible to take account of the bulk of the red corpuscles (see p. 30), a point of some value clinically.

When the *oligocythæmia* is very pronounced, a glance through the microscope will suffice to recognise it ; and so with *oligochromæmia*—diminution of haemoglobin—a little practice will enable us to detect it by a simple examination of the blood in a very thin layer without the addition of any fluid. To effect this, the end of the finger should be washed in plain water, and pricked, and the first drops of blood allowed to flow off. A glass slide should then be allowed to touch the summit of the drop of blood on the skin without coming in contact with the finger, quickly withdrawn, and a cover-glass placed over it.

In this way such impurities as epithelium, &c., are avoided.

The use of carbolic acid, æther, or alcohol, to wash the skin, is not to be recommended, since these bodies are likely to produce changes in the appearance of the corpuscles. When the object of the examination is the detection of micro-organisms in the blood, the utmost care must be taken in cleansing the skin (see p. 46).

Proceeding in this way in a case of oligocythæmia, when the slide is placed under the microscope, a marked diminution in the number of

blood-corpuses will be noticed. The red corpuscles will also in most cases be paler than normal; their usual bi-concave shape less marked; they are somewhat flattened, and they tend less to run into rouleaux or assume stellate forms. At the same time, they may be seen to have undergone peculiar changes of shape (*poikilocytosis*).

For many purposes it is well to fix the blood with some preserving fluid before examination. A solution of common salt (0.8–1.0 per cent.) or of sulphate of magnesium (5 per cent.) may be used (*Grüber*). *Hayem's* or *Pacini's* solution may be employed. *Hayem's* has been already described.



101

1

0.5

Pacini's solution is prepared thus:—A mixture is made of 1 part corrosive sublimate, 2 parts common salt, 13 parts glycerine, and 113 parts distilled water, and the fluid is allowed to stand for at least two months. When about to be used, a portion is diluted with three times its bulk of distilled water, and filtered through blotting-paper.⁴⁸

When it is a question of a lesser degree of oligocythaemia, this proceeding will not suffice, and we must have resort to special means of estimating the precise number of the corpuscles or the relative quantity of haemoglobin present. In recent times a great many instruments have been constructed with the first-named object—as those of *Quincke*, *Malassez*, *Hayem*, *Gowers*,⁴⁹ *Thoma-Zeiss*, and *Alferow*.⁵⁰ The principle on which all these are constructed is the same. A known quantity of blood is mixed in definite proportion with some indifferent fluid (3 per cent. salt solution, &c.), a portion of the mixture is placed upon a hollow slide of known contents and graduated surface, and then the corpuscles are counted with the aid of the microscope.

FIG. 2.—Capillary Tube (Thoma-Zeiss Apparatus for Counting the Blood-Corpuses).

(a.) **The Thoma-Zeiss Apparatus for Counting Blood-Corpuses.**—The simplest and best of these instruments is that of Thoma and Zeiss. It consists of a capillary tube of glass about 10 centimetres long, expanding in its upper third to a bulb, in which lies a small glass ball. The lower end of the tube is furnished with a scale, graduated in parts numbering 0.1, 0.5, 1, up to 101 (fig. 2). With this instrument is used a counting-chamber invented by *Abbe*⁵¹ and

(a.) The Thoma-Zeiss Apparatus for Counting

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Zeiss. This is a glass receptacle cemented upon a glass slide (fig. 3); it is exactly 0.1 mm. in depth, and its floor is marked out into microscopic squares (fig. 4). The space overlying each square = 1 4000 mm.³,* and the squares are portioned out in groups of 16 by planer lines (fig. 5). Similar counting cells have been devised by *Gabritschewsky*, *Zappert*, and *Elsholz*.

Application of the Process.—A puncture is made in the tip of the



FIG. 3



FIG. 4

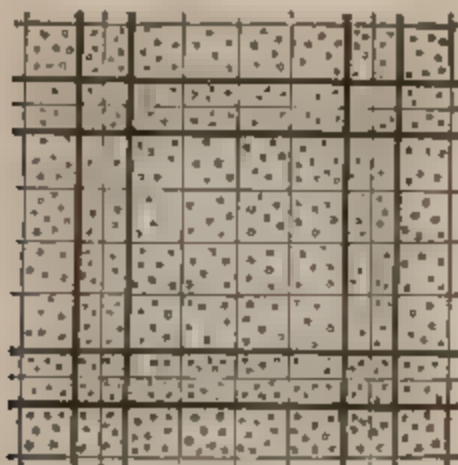


FIG. 5.—Thoma-Zeiss Apparatus for Counting Blood Corpuscles.

finger, and in the doing of this the precautions already indicated are taken. Blood from the summit of the exuding drop is then sucked into the tube until it reaches the mark .5 or 1. The point of the tube is wiped, and a 3 per cent. solution of common salt sucked in until the fluid has risen to the point marked 101. For some years the author has used Hayem's fluid in this experiment (see p. 7). *Daland*¹² and

* Mm.³—cubic millimetre.

*Sadler*⁵³ prefer a $2\frac{1}{2}$ per cent. solution of bichromate of potash. The contents of the tube are then thoroughly mixed, and the column of fluid in the capillary tube is removed by blowing into the tube, as the blood would not mix with the solution of common salt. To neglect this precaution would vitiate the experiment.⁵²

The capillary tube must be carefully cleaned after use, by washing it with distilled water, then with alcohol, and finally with æther, and a brisk current of air blown through it. For the latter purpose *Böhm's* air-pump is very suitable.

The hollow cell of the slide is next filled with the mixed blood-and-salt solution, care being taken to guard against the admission of air-bubbles, and the cover-glass is accurately adjusted in such a manner that Newton's colour rings are produced. The preparation is left to stand for some minutes, so as to allow of an intimate admixture of its parts, after which it is placed under the microscope, and looked at with a power of 30-70 diameters, when it will be seen whether any air-bubbles or foreign bodies are present in it, and whether the corpuscles are pretty evenly distributed through the fluid. The latter are then counted under a high power. In doing this, the number present in sixteen squares is counted, and from this the average is estimated. The greater the number of the squares taken, the more accurate will be the result attained. To count the corpuscles contained in sixteen squares, *Lyon* and *Thoma*⁵⁴ made the following suggestions:—

The space overlying a vertical series of four of the squares is taken as that the contents of which are to be estimated together. All the cells which cover or impinge upon the upper boundary of the rectangle formed by these four squares are to be reckoned, and that whether the cells themselves are situated within or without the boundary-line in question. So also are those which touch upon the line bounding the field of four-squares on one (left) of its sides, and all that are entirely included within its limits without being anywhere in contact with the boundary.

The object-glass used in this investigation should be *Zeiss*, C or D; *Hartnack*, 6; *Reichert*, 6; or *Gundlach*, V.

The estimation of the total number of corpuscles is conducted as follows:—If the blood in the tube reached to the point 0.5, its proportion in the mixed solution will be 1 : 200; if to the point 1.0, 1 : 100. Multiply the number of corpuscles counted in all the squares by 4000 ($\frac{1}{1000}$ being the cubic contents overlying a square), and the result by 100 or 200, according to the degree of dilution. Then divide the product by the number of squares taken, and the result gives the number of blood-corpuscles contained in a cubic millimetre of blood.

The following plan of making the calculation is employed in the

clinic at Prague:—The contents of five large squares are estimated by counting the corpuscles in the manner shown above, and with due precaution concerning the boundary-lines. Let a be the number of corpuscles estimated. Now, since one large square includes sixteen small ones, five large squares include eighty small ones, and their cubic content is $\frac{8}{1000}$ cubic mm., or $\frac{1}{125}$ cubic mm. Then 1 cubic mm. of the fluid contains $50a$; and the blood itself 5000 or 10,000 (according to its dilution) a of red corpuscles. *Miescher's* modifications of this apparatus include no improvements of any importance.⁵⁵

To estimate the number of *white* corpuscles in a specimen of blood, *Thoma*⁵⁶ dilutes the latter with water containing one-third per cent. glacial acetic acid in the proportion of 1:10. In this way the red corpuscles are destroyed, and the white alone remain in the field of vision. In mixing the fluids, the same observer employs a mixing-glass specially devised by Zeiss for the purpose. The process may also be carried out thus:—By means of a pipette of 1 cc. contents, and accurately graduated in 0.1 cc. units, 0.9 cc. of the acetic acid solution is measured out into a watch-glass; with another pipette holding exactly 0.1 cc., the blood is added to this, and the two well mixed. A drop of the mixture is placed within the counting-chamber of the cytometer prepared as before; and now, since the number of corpuscles is relatively fewer, the entire field, and not its marked-out divisions, is taken as the basis of calculation, greater accuracy being so obtained. To the same end, a lower power will be used, so as just to bring the marks in the floor of the chamber clearly into view. Before beginning to count the corpuscles, however, it will be well to focus with the fine adjustment of the microscope, and to make sure that the cells have all settled.

The cubic contents of that part of the chamber which corresponds to the field of vision can be ascertained in the following way:—The divisions of the chamber which appear in the field are first counted. Each of these measures $\frac{1}{25}$ mm. across (v. *supra*: the area = $\frac{1}{100}$ mm.², the cubic contents = $\frac{1}{1000}$ mm.³). The diameter of the field, therefore, is $\frac{1}{25}$ mm. multiplied by the number of divisions which it contains. Thus, for instance, if ten of the divisions are seen, the diameter of the field = $10 \times \frac{1}{25}$ mm. or $\frac{10}{25}$, and the radius = $\frac{10}{50}$ mm. The area of the field, therefore = $\pi (\frac{10}{50})^2$ mm.²; and if the chamber is 0.100 mm. in depth, its cubic contents = $0.1 \times (\frac{10}{50})^2 \pi$ cubic mm. Hence we obtain the following formula: *—

$$\frac{10 \times Z}{M \times Q}$$

where M = the number of divisions under the microscope, Z = the number of cells counted, Q = the cubic contents of the field (Q = 0.1 π

* $\pi = 3.1416$. mm.² = square millimetre. mm.³ = cubic millimetre.

R^2 , where R = the radius of the field in mm.), and where the blood is diluted in the proportion of 1 : 10. The formula will give us the number of cells contained in a cubic millimetre of undiluted blood. Where the degree of dilution is 1 in 10, and where, as is usually convenient, 16 squares are taken, from the general formula results the following :—

$$\frac{10,000 \times Z}{314};$$

or, if the degree of dilution be 1 in 20, the squares 16—

$$\frac{20,000 \times Z}{314}$$

i.e. the number of white corpuscles in 16 squares (Z) multiplied by 10,000 for the 1 in 10 solution, by 20,000 for the 1 in 20 solution, and divided by 314, gives the number of white corpuscles in a cubic millimetre of blood.

In the clinic at Prague it is the custom to count the corpuscles in the entire field of 400 small squares. Let this number be N. The cubic content of the field is $\frac{4}{1000}$ or $\frac{1}{10}$ cubic millimetre. Consequently, 1 cubic millimetre of the fluid contains 10 N corpuscles, and the blood, according to the degree of its dilution, 100 N or 200 N white corpuscles.

When there is a very great increase in the number of leucocytes, as in leukæmia, their number can be estimated in the same manner as that of the red corpuscles, and the relative proportion of the two can be determined at the same time with sufficient accuracy, if only an adequate number of squares is taken into account. Great assistance in such experiments may be derived from the use of a 3 per cent. salt solution coloured with gentian violet, in which the leucocytes are stained, and become readily discernible from the red blood-corpuscles, which are usually somewhat paler than normal. *Toison*⁵⁷ employs for the purpose a staining fluid of the following composition :—

Distilled water	160 cc.
Glycerine	30 cc.
Sulphate of soda	8 grms.
Chloride of sodium	1 grm.
Methyl-violet	0.025 grm.

For the same purpose, *Mayet*⁵⁸ recommends that the blood be mixed with peroxylic acid, glycerine, and water, which is said to confer upon the coloured blood-corpuscles a beautiful red tint, whilst the white corpuscles remain unaffected by it.

Zappert employs this fluid in the enumeration of eosinophil cells. According to *Marschner*,⁵⁹ Mayet's process has little value, and *Toison's*,

though useful in the enumeration of red corpuscles, offers no advantage in counting leucocytes. *Elzholz*⁶⁰ uses a dilution fluid composed of 7 parts of a 2 per cent. solution of eosin, 45 parts of glycerine, and 55 parts of water; with this, it is said, the enumeration of the leucocytes is greatly facilitated. Another process applicable to the purpose is that of *Müller-Rieder*.⁶¹ *Zappert*⁶² uses Mayet's fluid for estimating the eosinophil cells.

[*(b) Gowers' Hemacytometer.** This instrument is the most commonly used for clinical purposes in this country. "The haemacytometer (fig. 6) consists of—
(1.) A small pipette, which, when filled to the mark on the stem, holds exactly

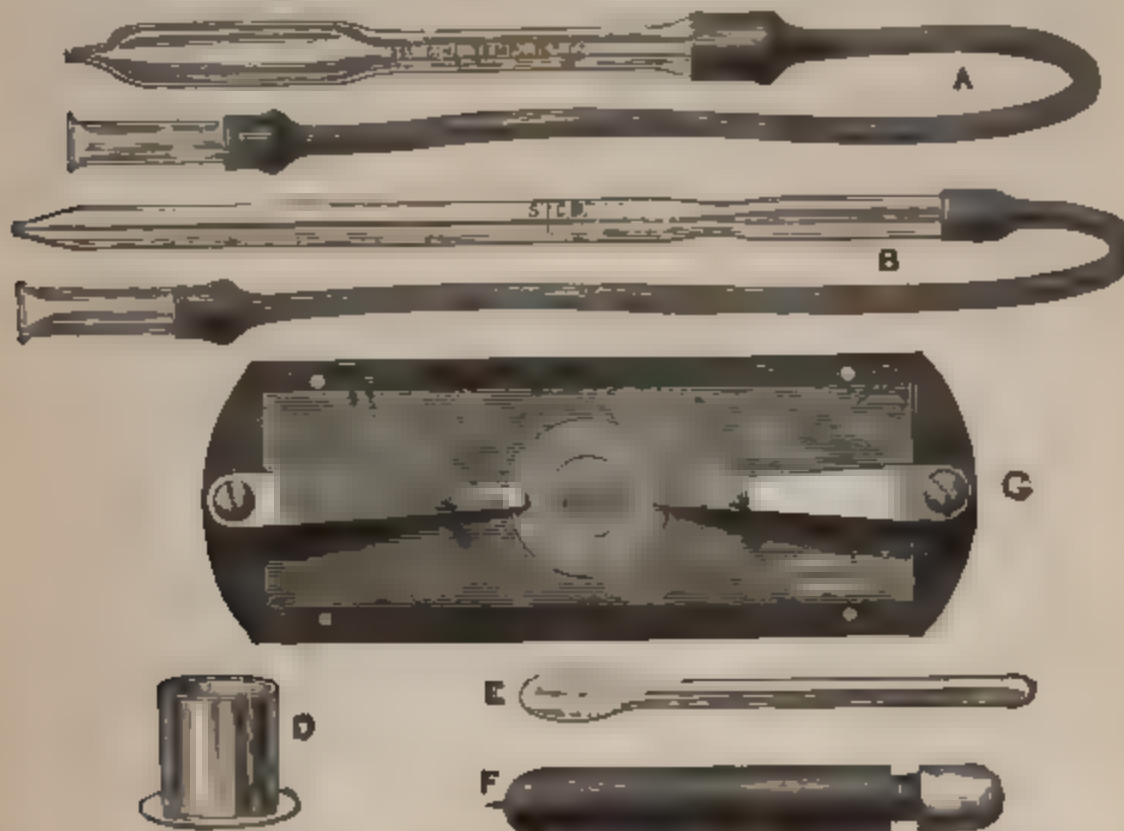


FIG. 6.—Gowers' Apparatus. A, pipette for measuring the diluting solution. B, capillary tube for measuring the blood. C, cells with divisions on the floor, mounted on a slide; D, vessel in which the dilution is made. E, glass-stirrer. F, guarded spear-pointed needle.

995 cubic millimetres. It is furnished with an india-rubber tube and mouthpiece to facilitate filling and emptying. (2.) A capillary tube marked to contain exactly 5 cubic millimetres, with india-rubber tube for filling, &c. (3.) A small glass jar in which the dilution is made. (4.) A glass stirrer for mixing the blood and solution in the glass jar. (5.) A brass stage-plate, carrying a glass slip, on which is a cell $\frac{1}{2}$ of a millimetre deep. The bottom of this is divided into

* For the description and figure we are indebted to *Landois and Stirling's "Physiology,"* vol. i. p. 5. *Fourth edition.*

[† In practice error is apt to arise from variation in the depth of the cell, and it is not easy to obtain one precisely $\frac{1}{2}$ mm. Assuming that the same instrument is always used by the observer, this is best corrected by ascertaining the true depth and allowing for the error in mixing the solution. Suppose, for instance, the cell is found to have a depth of $150\ \mu$ instead of $200\ \mu$, 5 parts of blood should be added to 945 of the diluting fluid, instead of to 995. The results so obtained will be absolutely accurate (*Hurt*).]

$\frac{1}{10}$ millimetre squares. Upon the top of the cell rests the cover-glass, which is kept in its place by the pressure of two springs proceeding from the ends of the stage-plate."

The method of employing the instrument is as follows :—The diluting solution used is a solution of sodic sulphate in distilled water, sp. gr. 1025, or the following :—Sodic sulphate, 104 grains ; acetic acid, 1 drachm ; distilled water, 4 oz. "995 cubic millimetres of the solution are placed in the mixing jar ; 5 cubic millimetres of blood are drawn into the capillary tube from the puncture in the finger and then blown into the solution. The two fluids are well mixed by rotating the stirrer between the thumb and finger, and a small drop of this dilution is placed in the centre of the cell, the covering-glass gently put upon the cell, and secured by the two springs, and the plate placed upon the stage of the microscope. The lens is then focussed for the squares. In a few minutes the corpuscles have sunk to the bottom of the cell, and are seen at rest on the squares. The number in ten squares is then counted, and this, multiplied by 10,000, gives the number in a cubic millimetre of blood."^{63]}

To estimate the COLOURLESS corpuscles only, the blood is mixed with ten parts of 0.5 per cent. solution of acetic acid, which destroys all the red corpuscles [Thoma].

The instruments of *Bizzozero*, *v. Fleischl*, *Hénocque* [and *Gouers*]⁶⁴ involve the second principle to which allusion has been made, viz., the estimation of the quantity of haemoglobin in the blood (see p. 9).

*Hedin's*⁶⁵ instrument for measuring corpuscles in bulk (see p. 28) is applicable to the same purpose.

Mention must be made of *v. Limbeck's*⁶⁶ researches on the subject of the resistance of the red corpuscles and the isotonic property of blood serum, since they may find a practical application by-and-by. The same potential importance attaches to *Laker's*⁶⁷ observations on the resistance of the blood-corpuscles. Neither of the foregoing methods has as yet any clinical bearing.⁶⁸

Professor *Wright*,⁶⁹ of Netley, has described a method of determining the state of blood coagulability under various circumstances. It requires half-a-dozen or a dozen capillary tubes of equal calibre and furnished with mouthpieces. The automatic pipette supplied with *v. Fleischl's* haemometer is suitable for the purpose. Into these pipettes at regular intervals (which must be carefully noted) is drawn up a drop of blood, which is allowed to remain in the tube. In about two minutes from the time of its filling, the blood in the first tube is examined, and afterwards that in the others at intervals of one to one and a half minutes. This is done by blowing through the tubes to see if they are yet blocked, and receiving the contents when removed upon clean blotting-paper, so that it may be seen whether they are fluid or solid. Variation in the time of clotting can thus be shown to occur in different conditions. *Wright* maintains that the blood coagulated more quickly after the administration of calcium chloride. The author cannot confirm this statement ; nor can he, as the result of observations in a great many cases of haemophilia, vouch for any therapeutic virtue as belonging to calcium chloride.

(c.) **Bizzozero's Chromo-Cytometer.** [According to *Bizzozero*⁷⁰ a knowledge of the number of coloured blood-corpuscles is of less practical value than to know the quantity of haemoglobin, and, as a matter of fact, the amount of the latter is not necessarily in direct proportion to

the number of the former. For the purpose of estimating the *amount* of haemoglobin *Bizzozero* has invented a small, practical, and handy instrument which he calls a *chromo-cytometer*⁷¹ (fig. 7).

By means of this instrument we can estimate the amount of haemoglobin in the blood, and it can be used either as a cytometer or as a chromometer. In both cases it is essentially an expedient for varying the thickness of a stratum of blood.

To use it as a cytometer, the blood is mixed with a definite volume (1 : 50) of an indifferent solution, e.g. normal saline solution (0.75 grammes of sodic chloride in 100 cc. water), so that the corpuscles remain intact in the fluid. The quantity of haemoglobin is estimated by the thickness of a layer of fluid through which one in a dark room can distinctly see the edges of a candle-flame placed at a distance of 1½ metres from the instrument.

In using the instrument as a chromometer, the blood is mixed with a known volume of water, which dissolves out the haemoglobin from the corpuscles. The amount of haemoglobin is then calculated from the thickness of the layer of this mixture, which yields a covering exactly equal in intensity to that of a red-coloured glass supplied with the instrument.

The chief part of the instrument consists of two tubes (*ab*, *cd*), working one within the other, and closed at the same end by glass discs (figs. 7 and 8), while the other ends are open. The one tube can be completely screwed into the other, so that both glasses touch. Connected with the outer tube is a small open reservoir (*r*), from which fluid can pass into the variable space between the two glass plates at the ends of the tubes. By rotating the inner tube, the space between the two glass plates can be increased or diminished, on the principle of Hermanu's haematoscope, and the screw is so graduated as to indicate the distance between the two plates, i.e. the thickness of the layer of fluid between them. Each complete turn of the screw = 0.5 mm., and the subdivisions on it are so marked—25 to one turn (index, fig. 7, *cd*)—that each subdivision of the index = $\frac{0.5}{25} = 0.02$ mm. When the inner tube

is screwed home and touches the glass disc in the outer tube, the index stands at 0 on the scale. If the instrument is to be used merely as a cytometer, these parts suffice; but if it is to serve as a chromometer, the coloured glass is needed also. The instrument is also provided with small glass thimbles with flat bottoms, containing 2 and 4 cc. respectively; a pipette graduated to hold ½ and 1 cc., and another pipette for 10 and 20 cmm., the latter provided with an india-rubber tube, to enable the fluid to be sucked up readily; a bottle to hold the saline solution, and a glass stirrer.

To Use the Instrument as a Cytometer.—1. By means of the pipette place .5 cc. of normal saline solution in a glass thimble.

2. With a lancet or needle puncture the skin of the finger at the edge of the nail.

3. With the pipette suck up exactly 10 cmm. of blood, observing the precautions already indicated at p. 9. Mix this blood with the .5 cc. saline solution, and suck part of the latter several times into the capillary tube, so as to remove every trace of blood from the pipette. Mix the fluids thoroughly. Carefully cleanse the pipette with water.

4. Pour the mixture into the reservoir (*r*) of the instrument. Gradually rotate the inner tube, and as the two glass discs separate, the fluid passes into the space between them.

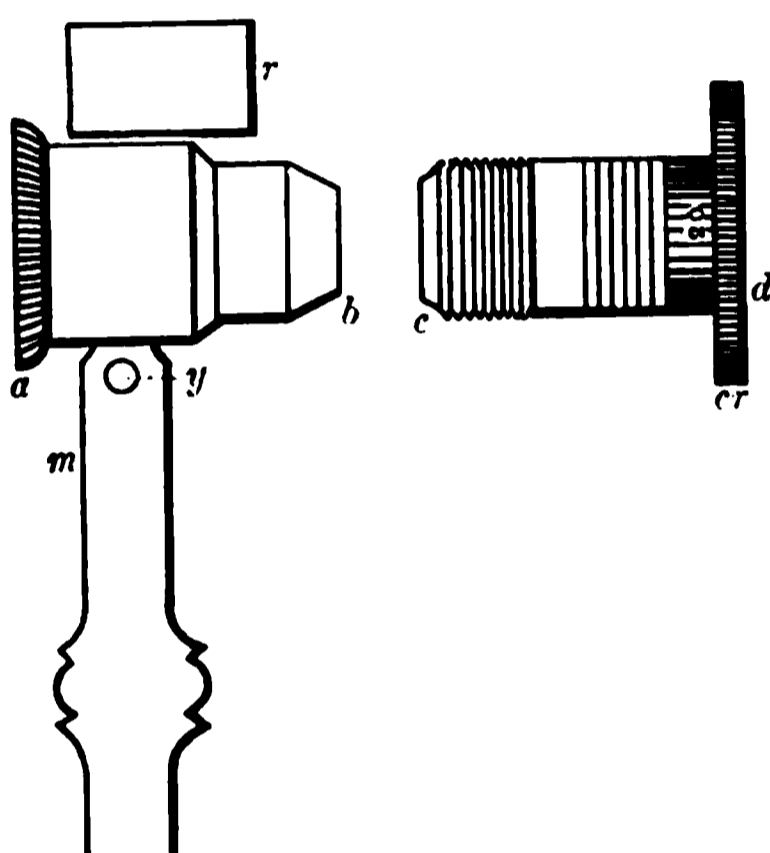


FIG. 7.—General view of the instrument. *ab* and *cd*. Two tubes, the one fits inside the other; *r*. Reservoir communicating with the space between *c* and *b* when *cd* is screwed into *ab*; *cr*. Milled head, and index-scale to the left of it; *y* for *ast* of fig. 9; *m*. Handle.

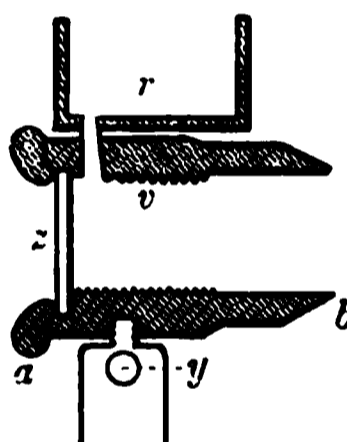
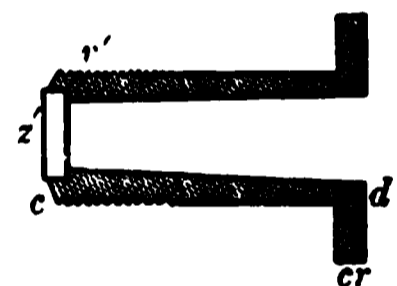


FIG. 8.—Showing how *cd* fits into *ab*. *z* and *z'*. Plates of glass closing the ends of *ab* and *cd*; other letters as in fig. 7.



5. In a dark room light a stearin candle, place it at a distance of $1\frac{1}{2}$ metres, and, taking the instrument in the left hand, bring the open end of the tubes to the right eye. With the right hand rotate the inner tube to vary the thickness of the column of fluid, and so adjust it until the outlines of the upper three-fourths of the flame can be distinctly seen through the stratum of fluid. Vary the position of the inner screw so as to determine accurately when this occurs. Read off on the scale the thickness of the stratum of fluid.

Graduation of the Instrument as a Cytometer.—In this instrument the graduation is obtained from the thickness of the layer of blood itself, and the amount of haemoglobin is calculated directly from the thickness of the layer of blood which is necessary to obtain a certain

optical effect, viz., through the layer of blood corpuscles to see the outlines of a candle-flame placed at a certain distance.

From a number of investigations it appears that in healthy blood the outlines of the flame of a candle are distinctly seen through a layer of the mixture of blood $\frac{110}{100}$ mm. in thickness.

Let the number 110 correspond to 1, or, better still, 100 parts of haemoglobin; then it is easy to calculate the relative value of the subdivisions of the scale on the tube of the instrument. Let g = the degree of the scale for normal blood; g' , that for the blood being investigated; e , amount of haemoglobin in the former; and e' , the amount sought for in the latter.

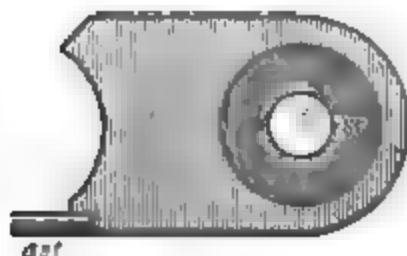


FIG. 9.—Coloured glass, *f*, in a blackened brass screen, *l*; *ast* Stem for fixing it in *y* of fig. 7; *sc*. Brass tube in which *f* is fixed; this is used when the instrument is employed as a chromometer.

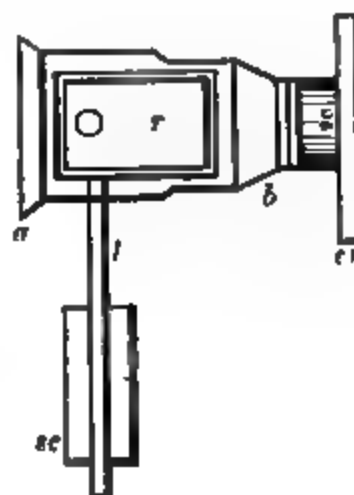


FIG. 10.—Instrument seen from above. Letters as in other figs. (7, 8, 9).

Assuming that the product of the quantity of haemoglobin and the thickness of the stratum of blood is constant, so that

$$e g = e' g'.$$

Then

$$e' = \frac{eg}{g'}.$$

Let us assume that the blood investigated gave the number 180; then, using the above data:—

$$e' = \frac{100 \cdot 110}{180} = \frac{11,000}{180} = 61.1.$$

The blood, therefore, contains 61.1 parts of haemoglobin. The following table gives the proportion of haemoglobin, the normal amount of haemoglobin being taken as = 100:—

Cytometer Scale.	Haemoglobin.	Cytometer Scale.	Haemoglobin.
110	100.0	170	64.7
120	91.6	180	61.1
130	84.6	190	57.9
140	78.5	200	55.0
150	73.3	210	52.4
160	68.7	220	50.0

If the instrument be used as a chromometer, the blood is mixed with a known volume of water, whereby the haemoglobin is dissolved out of the red corpuscles and the fluid becomes transparent. The quantity of haemoglobin is calculated from the thickness of the stratum of fluid required to correspond exactly to the colour-intensity of a tinted-glass which is attached to the instrument. The colour-intensity of the glass is that of a definite solution of haemoglobin (fig. 9, f).

To use the Instrument as a Chromometer.—1. Place the coloured glass with its brass frame in the instrument (*ast* of fig. 9 in *y* of fig. 7).

2. With the necessary precautions (p. 9) mix 10 cmm. blood with .5 cc. distilled water, whereby in a few seconds a transparent solution of haemoglobin is obtained.

3. Pour this solution into the reservoir (*r*), and rotate the inner tube so that the fluid passes between the two glasses. Direct the instrument towards a white light or the sky, not towards the sun, and compare the colour of the solution with the standard coloured glass, a proceeding which is facilitated by placing a milky glass between the source of light and the layer of blood so as to obtain diffuse white light. When the two colours appear to have as nearly as possible the same intensity, read off on the scale the thickness of the layer of blood, and from this by means of the accompanying table ascertain the corresponding amount of haemoglobin.

This is done in the same way as for the cytometer, but the graduation is different, as in the one case we have to do with a candle-flame, and in the other with a coloured glass.

In very pronounced cases of anaemia, even with a layer of blood 6 mm. in thickness—the limit for which the instrument is constructed—the intensity of the mixture of blood may be less than that of the coloured glass. In such a case, instead of 10 cmm. of blood, 20 cmm. should be used.

Graduation of the Chromometer.—As the coloured glass has not absolutely the same intensity of colour in all chromometers, one must first of all estimate the colour-intensity of the glass itself. This is most easily done by ascertaining in a given specimen of blood what degree of the chromometer corresponds to the scale of the cytometer of the same blood.

Suppose that a specimen of blood by means of the cytometer gave 110, and by the chromometer 140; the number 110 of the cytometer = 100 haemoglobin, so that the chromometer number 140 must also be = 100. With the aid of the formula (p. 19) a similar table can be constructed for the chromometer. Suppose the blood investigated = 280; then by the aid of the formula and the data from normal blood:—

$$c' = \frac{100 \cdot 140}{280} = \frac{14,000}{280} = 50.$$

This blood, therefore, contains 50 parts of haemoglobin.

Example.—Blood gives 130 with the cytometer and 190 with the chromometer; what is the initial number of the chromometer graduation corresponding to 100 parts of haemoglobin?

If 130 (cytometer) corresponds to 190 (chromometer), then 110 cytometer (*i.e.* graduation corresponding to 100 parts of haemoglobin) corresponds to x chromometer graduation:

$$130 : 190 = 110 : x \therefore x = \frac{190 \cdot 110}{130} = \frac{20,900}{130} = 160.7.$$

Blood containing 100 parts haemoglobin will correspond to 160 of the chromometer scale, and beginning with this number as a basis, with the aid of our formula it is easy to construct a table showing the relation.

Whilst the value of the cytometer scale remains the same for every instrument, the chromometer scale varies with each instrument, as the colour-intensity of the glass is not necessarily the same in all. But it is easy to construct a scale for each instrument by investigating a specimen of blood and comparing it with the cytometer graduation as indicated in the foregoing paragraph.

In using the instrument certain precautions must be observed. The exact quantity of the several fluids must be carefully measured; evaporation must be prevented by covering the blood-mixture. Further, it is well not to look at the fluid too long at a time, as the eye becomes rapidly fatigued.

In cases of leukæmia, where there is a large number of white corpuscles rendering the mixed fluid opaque, the corpuscles may be made to disappear by adding a drop of very dilute caustic potash. If the opacity does not disappear by the addition of this substance, then the opacity is due to the presence of fatty granules in the blood, so that by this means we can distinguish lipæmia from leukæmia.

Further, the operation must not be carried out too slowly, as the saline solution only retards the coagulation of the blood and does not arrest it.

Bizzozero claims that when the instrument is used as a cytometer the mean error is not greater than 0.3 per cent.] *

Sadler,⁷² working in the author's clinic, has found that *Bizzozero's* instrument gives very accurate results.

*Oertel*⁷³ has applied it in a very scientific manner to determine the "coefficient of density" of the blood. It is necessary to mention here *de Thierry's* hæmospectroscope⁷⁴ as an instrument of which the author and the editor have as yet no personal experience.

* The instrument may be obtained from F. Koristka, Via Circo, 14, Milan, and costs thirty-five lire.

(d.) **v. Fleischl's Hæmometer.***—The application of this instrument (fig. 11) depends upon the principle that the colour of the blood diluted with water may be compared with that of a glass wedge tinted with Cassius's golden-purple, or some such pigment.

Its essential part is the red glass wedge, which is mounted movably beneath a platform like that of a microscope, with a circular opening in its centre. Upon this the light from a gas or oil lamp (daylight is not



FIG. 11.—Von Fleischl's Hæmometer.

admissible) is projected by a plate of plaster of Paris. Above the wedge, and exactly over the circular opening in the platform, is fixed a metallic tube 1½ cm. long, closed at the bottom with a plate of glass, and divided by a vertical metallic partition, so that one-half of the metallic tube receives its light through the red glass wedge, the other directly from the white reflector. When the apparatus is in use, the former of

* Von Fleischl's instrument is made by Reichert of Vienna, and sold for thirty-five florins.

these compartments is filled with pure water, the other with water mixed with a known quantity of blood.

To secure this known quantity, *von Fleischl* has designed an automatic blood pipette of such a capacity that, when healthy blood is used, the resulting mixture corresponds in colour to that derived from the part of the red glass wedge which is marked 100. From this point to its sharp edge (where 0 stands), the wedge is graduated in ten divisions, which represent its diminishing thickness, the Nos. 90, 80, &c., being marked on the apparatus.

The instrument is employed thus:—The blood is obtained from a puncture in the finger, and placed by means of the pipette in the proper compartment of the tube. Both compartments are then filled with water, and the red glass wedge is moved until the two fluids show an equal intensity of red colour. The number indicated on the scale is then read off. Suppose this should be 80,—then the blood examined contains but 80 per cent. of the normal proportion of hæmoglobin, or the quantity of hæmoglobin is to that of healthy blood as 80 : 100. Now, assuming that in a healthy man 14 grms.* may be taken as the amount of hæmoglobin in 100 grms. of blood, we can calculate the latter absolutely for the specimen examined by means of the formula :

$$X = \frac{14 \times R}{100}$$

Where

X = the quantity of hæmoglobin in 100 grms. of blood :

R = the figure obtained with von Fleischl's apparatus to express the relative proportion of hæmoglobin in the blood ; and

14 = the normal quantity of hæmoglobin in the blood of a healthy adult.†

Although it must be conceded that the results obtained with this instrument are not absolutely correct, it still supplies a simple and ready means of estimating the hæmoglobin in the blood, and has the further advantage that it needs but little of the latter to work upon. It has proved a useful adjunct to the *Thoma-Zeiss* apparatus, or that of *Gowers*, in the investigation of changes in the blood for clinical purposes.

As an instrument for the estimation of hæmoglobin, its utility is vouched for by a great number of observers (*Gottlieb, Laker, Barbacci, Kisch, J. Meyer, Hæberlin, Widowitz, Stierlin, Schiff, Wilkens, Reinl*).⁷⁵ The hæmometer is said by *Lederer*⁷⁶ to be in no way inferior to *Gowers'*

* *J. G. Otto* states the normal quantity at 13.77 per cent. 14 is taken here (as by *Hénocque* elsewhere) as a simpler expression for the purposes of the calculation.

† This number is chosen here so as to afford a basis of comparison between the results obtained with von Fleischl's instrument and those with *Hénocque's*, which will be described presently.

apparatus, which is more extensively used in England, but it is necessary to recognise that, with either, the results obtained are only approximate. Greater accuracy may be had by the spectro-photometric processes, employed by *von Renl* amongst others. The so-called improvements of *von Fleischl's* instrument devised by *Miescher*⁷⁷ are of small account.

(e.) **Hénocque's Hæmatoscope.**⁷⁸—This instrument has the advantage over others employed for a similar purpose, that while comparatively little fluid is needed for its application, this consists of pure blood undiluted with artificial serums.

It consists essentially of two glass plates superimposed in such a manner as to enclose a prismatic capillary space. The inferior of these plates is the broader. Upon the upper part of its surface is engraved a millimetre scale, 0 to 60, reading from left to right, and at either ex-



FIG. 12. Section of Hæmatoscope



FIG. 13. Hénocque's Hæmatoscope filled with Blood.

tremity it carries a cap of nickelled metal, in which is a groove for the reception of one end of the smaller (upper) plate. These grooves are so placed that whilst the plates are in immediate contact at one end, that opposite 0 of the scale, they are separated at the other by an interval of 0.3 mm. The smaller plate can be made to slide in the grooves under gentle pressure, and can thus be removed for the purpose of cleaning the instrument. From the above description it will be seen that a layer of blood introduced within the capillary chamber has a thickness varying uniformly from 0 on the left to 0.3 mm. under the mark 60 on the graduated scale; and it also follows that its depth increases by 0.005 mm. for every mm. distance towards the right. Thus its depth at any point may be known by multiplying the corresponding figure on the scale by five, when the result is expressed in thousands of a millimetre or micron.

The prismatic chamber is filled with blood obtained by pricking the finger, and this is done best by bringing the lower lamina on a level with the puncture, and permitting the issuing drops to flow upon it at a gentle decline. The blood will then arrange itself in an even layer; and if its continuity be interrupted by empty spaces or air-bubbles, these can be extruded by slightly tapping the glass wall with the finger-nail. Six drops generally suffice. The edges of the instrument, when filled, are wiped clean, and the examination may be begun.

This is conducted in two different ways:—

(a.) The first (*procédé diaphanométrique*), the readier and simpler, has for its object to estimate the relative opacity of the blood, and so to infer the proportion of hæmoglobin which it contains.

For this purpose there is supplied with the hæmatoscope a plate of white enamelled metal, bearing on its upper part a millimetre scale precisely similar to that engraved upon the lower of the two glass plates, and below, a descending series of figures, of which the first underlies the

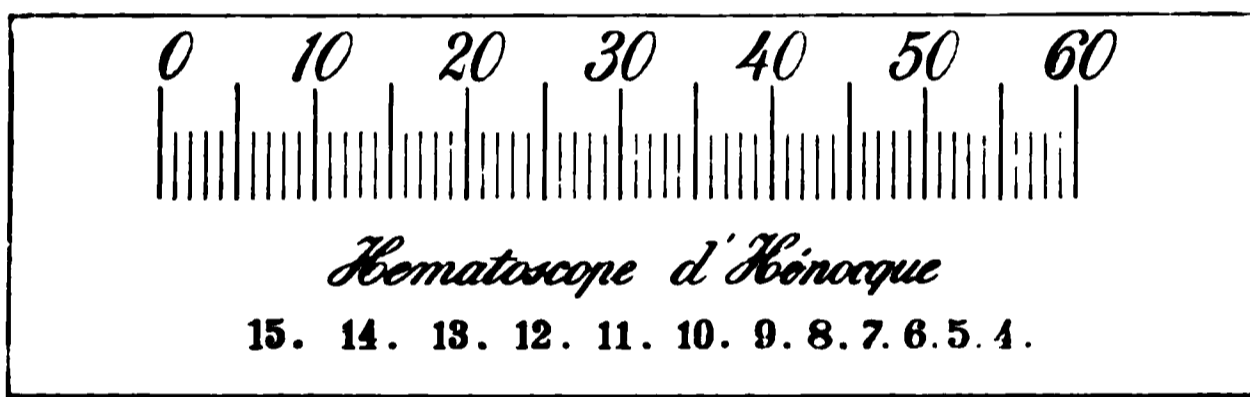


FIG. 14.—Enamelled Plate belonging to the Hæmatoscope.

8 mm. mark above. Then follow at constantly diminishing intervals the figures 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4. The markings upon this plate are traced throughout in black. When in use, it is placed behind the hæmatoscope filled with blood, and held there in such a position that like markings upon the millimetre scales accurately correspond. It is evident that the portion of the blood layer which is thinner, and therefore less deeply coloured, will be transparent, and will suffer the marks beneath to be visible, while these disappear towards the thicker end. The examination consists in noting the point at which the figures (15–4) in the lower series cease to be visible; and it is clear that this point will be attained the sooner according as the blood is richer in hæmoglobin.

Further, Hénocque has arranged the series so that the second figure (14) expresses in grms. the quantity of oxyhæmoglobin in 100 grms. of blood, and this figure terminates the series as seen through a layer of blood of normal constitution. With blood taken from a case of anæmia on the other hand, the figure 8 or 7 may be legible, and this implies

that such blood contains in 100 grms. only 8 or 7 grms. of oxyhaemoglobin. Finally, the thickness of the stratum of blood at the point of requisite opacity may be ascertained from the mm. scale in the manner already indicated.*

*Hellström, Loos,*⁷⁹ and the author have satisfied themselves that the results obtained in this way are not to be relied upon, and that the figures generally indicate too high a proportion of oxyhaemoglobin.

(β.) In the second and more accurate mode of using the haematoscope the enamelled plate is dispensed with, and a Browning's spectroscope is required. The instrument, filled with blood as before, is placed opposite the cleft of the spectroscope, and the point is observed at which the characteristic spectrum of oxyhaemoglobin is first distinctly formed,—when the corresponding point on the millimetre scale of the glass plate is read off. The less haemoglobin in the blood, the thicker must be the layer from which a spectrum is obtained. In order to secure a correct reading from the scale, it is well to place the apparatus holding a stratum of blood upon a sheet of white paper against a window so as to examine it by bright and diffused daylight, and then directing the spectroscope over 1 or 2 cm. of its surface, the observer should several times judge for himself concerning the point of earliest definition of the spectrum. Of the numbers obtained in this way (which will usually differ by an amount expressing only two or three millimetres) the mean is taken, and employed for the purpose of the calculation. It must be allowed that the conclusion in this respect is always somewhat arbitrary, and leaves room for a difference of opinion as to when precisely the spectrum is formed; but once the eye has become accustomed to look for a certain clearness in the outline of the bands, it seeks for and easily appreciates it in every instance.†

From the reading on the scale at the point where the spectrum is thus seen, the thickness of the blood stratum, and the quantity of oxyhaemoglobin in a known quantity of blood, can be readily determined. In the case of normal blood, which contains 14 grms. of oxyhaemoglobin in 100 grms. of the fluid, the absorption-bands are plainly visible in the situation of the figure 14 on the mm. scale; and from what has been already said it follows that the thickness of the blood stratum at this point is 14×0.005 mm. = 0.07 mm. Let it be assumed now that

* [Hénocque's haematoscope may be obtained from M. Lutz, 82 Boulevard Saint-Germain, Paris. The price is twelve francs, and the enamelled plate costs five francs additional.—(ED.)]

† [This difficulty may be further obviated by the use of Hénocque's double spectroscope, by means of which two persons are enabled to make the observation at once. For a description of this instrument the reader is referred to the original communication, "L'Hémato-spectroscope." Compt. Rend., Soc. de Biologie, October 1886.]

in a given case the bands just become distinctly evident at a point corresponding to the division 20 on the index ; then the thickness of the layer which yields them is $20 \times 0.005 = 0.1$ mm. From these data the quantity of oxyhæmoglobin in 100 grms. of the blood may be calculated by the following equation :—

$$x : 14 = 0.07 : 0.005y$$

$$x = \frac{14 \times 0.07}{0.005y}$$

x = the quantity of oxyhæmoglobin sought.

In this formula :

14 = the quantity of oxyhæmoglobin in 100 grms. of healthy blood.

0.07 = the thickness of the blood stratum, which will make the absorption-bands plainly evident where the blood, as in normal blood, holds 14 grms. oxyhæmoglobin in 100 grms. of the fluid.

0.005 = the thickness of blood stratum corresponding to 1 mm.

y = the number of mm. read off at the point where the absorption-bands become distinctly visible. From this results the simple expression :

$$x = \frac{14 \times 0.07}{0.005y} = \frac{196}{y}$$

In the example chosen :

$$y = 20 \text{ and } \frac{196}{20} = 9.8,$$

i.e. the blood investigated contained 9.8 grms. of hæmoglobin in 100 grms.

To obviate the necessity for making the calculation afresh in each case, Hénocque has compiled a table from which the quantity of hæmoglobin may be deduced directly from the depth of the blood stratum.

Comparisons which the author has instituted between the results arrived at in this way with others derived by means of v. Fleischl's apparatus, have shown that the two are sufficiently in accord. Henschel,⁸⁰ however, is of opinion that the latter are more accurate, and accounts for this by pointing out that they are obtained from oxyhæmoglobin, whereas with Hénocque's hæmatoscope that body is still within the cells.

The preference will usually be given to v. Fleischl's apparatus, because of the greater quantity of blood needed for the application of Hénocque's hæmatoscope ; but the latter is especially suitable for the observation of such changes in the blood as the formation of methæmoglobin, &c., which can be recognised by spectrum-analysis. To detect such changes, Hénocque⁸¹ has applied it in a very ingenious manner. He observed the development of oxyhæmoglobin-bands in transparent parts, such as the lobe of the ear and the ungual phalanges of the finger, capable of

being illuminated by diffuse sunlight.* Then, in the case of the unguial phalanx, the part was ligatured, and he noted the length of time required for the appearance of the broad absorption-band of reduced haemoglobin. Proceeding in this way, he found that with a normal proportion of oxyhaemoglobin, reduction took place in the course of 70 seconds, while with anaemic blood the interval was shortened to 30-40 seconds.

As a result of his researches, *Hénocque* has arrived at the following formula, which is applicable to clinical purposes :—

$$E = \frac{M}{D} \times 5,$$

where

E = the energy of reduction ;

M = the mean proportion of haemoglobin ascertained by his method ;

D = the time in seconds in which reduction is accomplished.

The formula is derived from the following considerations :—In a specimen of blood holding 14 grms. oxyhaemoglobin in 100 grms., reduction takes place in 70 seconds, and in another holding 13 grms. in 100, it takes place in 65 seconds. Inspection of the figures shows that in both cases a fifth part of the quantity of oxyhaemoglobin (in 100 grms.) is reduced. Hence, to obtain the value of E (energy of reduction), the quantity of oxyhaemoglobin found is multiplied by 5, and the product divided by the number expressing the time (in seconds) in which reduction takes place. The apparatus may be employed also for the examination of milk, in the spectrum-analysis of urine and morbid fluids, and for the aniline dyes of so much consequence in staining processes. It seems to merit a description here on account of its extended utility ; and in any case, in connection with the spectroscopic examination of the blood, it must be classed with the apparatus of Thoma-Zeiss, Gowers, and v. Fleischl as a valuable addition to our resources.

(f.) **Hedin's Hæmatocrit.**^{s2}—This instrument is of great value. It enables one readily to estimate the volume of the red blood-corpuscles. Its parts are : (1.) A capillary tube for measuring and mixing the blood. Hedin uses a special contrivance for this purpose, but the mixing-glass noticed on p. 13 for counting leucocytes serves equally well. To prevent coagulation Hedin sucks Müller's fluid into the tube and then blood in equal quantities. These are then expelled into a small platinum dish and well mixed. *Daland*,^{s3} in the author's clinic, has found that a 2.5 per cent. solution of bichromate of potash answers best.

(2.) Two glass tubes, 35 mm. long, with a lumen 1 mm. in diameter and graduated into 50 parts.

(3.) A metallic frame, terminating at either side in an angle to support a small cylindrical recess, of the same diameter as the tubes (2.), and covered with a caoutchouc disc. From the middle of this metallic frame there projects downwards a hollow metal cylinder, by means of which it

* [In researches of this character the double spectroscope is specially recommended by *Hénocque*.]

can be made to rotate on a vertical axis (fig. 15). Connected with the central cylinder are two metallic springs, placed opposite each other, and carrying each at their upper ends, on a level with the recesses mentioned above, a caoutchouc cap. Between the recess and this caoutchouc cap on either side is placed one of the glass tubes which has previously been filled with the mixture of blood and Muller's fluid (or bichromate solution). In this position the tubes are closed by the caoutchouc cap, which is held against them by the pressure of the springs.

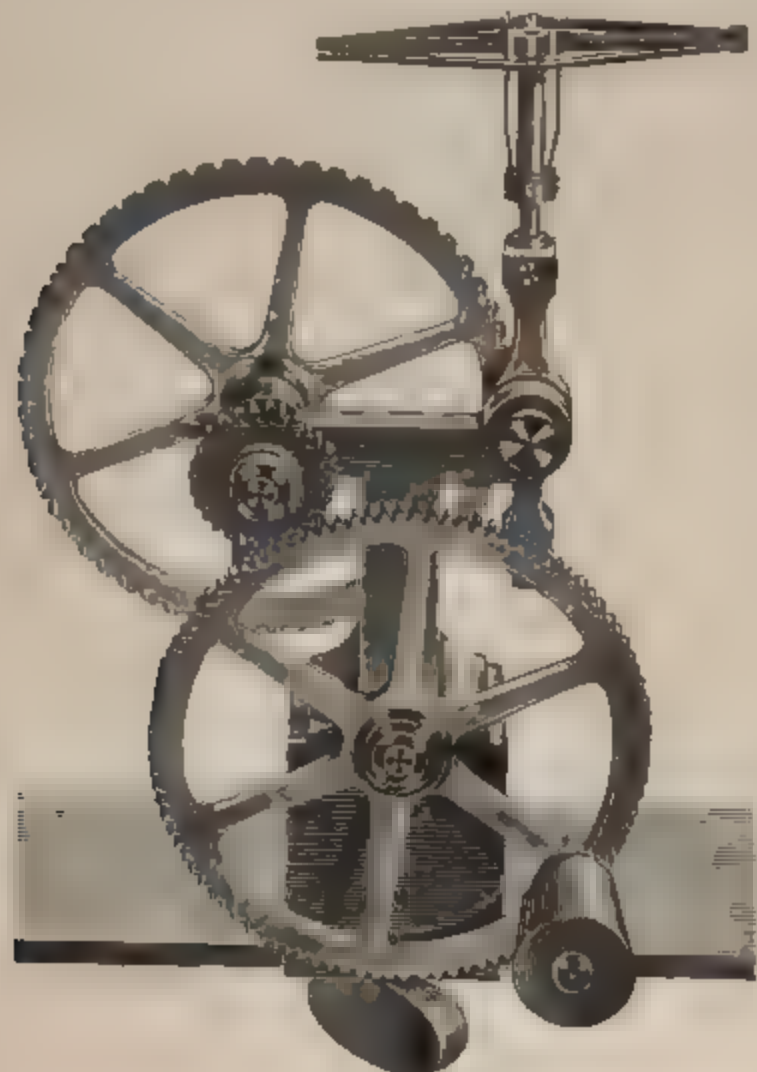


FIG. 15.—Hedin's Hematocrit.

(4.) A vertical support which can be made to rotate.

The instrument is used thus: With *Hedin's* capillary tube, or the mixer for estimating leucocytes, a mixture is made of blood, and a 2.5 per cent. solution of bichromate of potash, and with this the tubes are filled by plunging one end in the fluid, which is then drawn into the tube by the mouth applied to a piece of india-rubber tubing attached to the other end. The tubes being filled in this way are placed in position in the frame—the extremity towards the recess of the frame being first adjusted, and the caoutchouc caps of the springs being then made to

press on the other extremity. The frame is then attached by its vertical cylinder to the body of the instrument and made to rotate rapidly. The red corpuscles, in virtue of the centrifugal action, separate from the leucocytes and serum. After 50-70 seconds (where the bichromate solution is used) the volume of the layer of red corpuscles remains constant. The contents of the tubes are then arranged in three parts; at the distal end are collected the red corpuscles, forming a dense dark-coloured mass; next in order a small turbid band, consisting of leucocytes, and in health of a whitish colour; finally, the clear serum, which is coloured a bright yellow by Müller's fluid. To guard against error, a sheet of white paper is placed behind the tubes, and the volume of the red corpuscles is read off on the scale. The corresponding number multiplied by 4 gives the volume of red corpuscles in 100 parts of blood, as will appear from the following considerations. The volume of red corpuscles as read upon the scale is that contained in a mixture of equal parts of blood and bichromate solution, forming a column 35 mm. long and divided into 50 equal parts. The proportion of corpuscles in pure blood would be twice as great, and in 100 parts (instead of 50) again twice as great; therefore the percentage bulk may be expressed by the figure read off the scale multiplied by 4.

The description given here applies to an instrument furnished by *Sendling Sandström* of Lund, in Sweden, and differs somewhat from that published by *Hedin* himself.⁸⁴

This method is very serviceable in discriminating between the various diseases which affect the blood, and it may be used in part to replace the more difficult processes for counting the corpuscles. It may be substituted for these in conditions where the relative bulk of the red corpuscles depends only upon their number, and not also (as in many diseases, e.g. pernicious anaemia) upon their size. As to how far the results obtained in this way tally with those arrived at by counting, and under what circumstances the one method may supersede the other, the reader should consult *Daland*'s notice. In the same way it is possible to estimate approximately the relative proportion of red corpuscles and leucocytes, as, for instance, in certain cases of persistent leucocytosis. Observations made on numerous cases of leukæmia, have satisfied the author that the method suffices to determine the defective state of the blood in that disease. Further, it can be applied to the purpose of studying the leucocytes in the blood, and for the detection of micro-organisms therein.⁸⁵ The hæmatocrit does not answer the requirements of exact analysis; the author concurs with *Bleibtreu*'s⁸⁶ criticism on this head. It is nevertheless for many purposes a useful and practical instrument.

2. Leucocytosis.—By this term is meant a condition in which the number of white corpuscles in the blood is temporarily much increased. Such an increase occurs in health as part of the process of digestion.⁸⁷ One to two hours after the principal meal the white corpuscles are found in the blood in the proportion of 1 : 150 or even 1 : 100 of red corpuscles, diminishing soon after to 1 : 350 or 600, or, according to *Gräber*,⁸⁸ between 1 : 521 and 821. A great number of observations, conducted in the author's clinic, give the proportion to leucocytes of red corpuscles as 1 : 500–800⁸⁹ in healthy adults. *Reinecke's*⁹⁰ figures are 1 : 720.⁹¹ In new-born children the condition is different (*Schiff*).⁹² In the first three days of life the number of white corpuscles is very great, and after that they diminish, as also do the red corpuscles, till the proportion is 1 : 188—1 : 168. Much greater degrees of leucocytosis, which for the most part is transitory, are found in disease. *Virchow*⁹³ maintains that every affection in which the lymphatic glands are involved leads to this condition. It invariably accompanies croupous pneumonia (*Tumias*).⁹⁴ The author has never failed to find it in the croupous pneumonia of children. *V. Limbeck*, *Pick*, and *Lahr* confirm this. The first-named observes that it always follows exudation processes, and he proposes to give this form the name of inflammatory leucocytosis. [V. *Jaksch*⁹⁵ has quite lately observed that the prognosis in cases of acute pneumonia running its course without an increase in the number of white blood cells is very unfavourable.] *Sobotka*⁹⁶ has shown that leucocytosis follows vaccination, and that notable changes in the number of white corpuscles occur in the prodromal stages of measles, scarlet fever, small-pox, chicken-pox, and pneumonia. Leucocytosis does not arise from typhoid (*v. Limbeck*),⁹⁷ but may occur in its course. When it does so, it seems to point to a complication with some suppurative disorder (*Sauller*).⁹⁸ The condition occurs in connection with certain tumours,—sarcoma and cancer of rapid growth (*Sadler*, *v. Jaksch*),⁹⁹ in pernicious anaemia and chlorosis, regularly in the reaction stage resulting from the injection of Koch's fluid,¹⁰⁰ and in epidemic cerebro-spinal meningitis.¹⁰⁰ Leucocytosis is commonly an accompaniment of septic processes (*v. Jaksch*, *Rieder*),¹⁰¹ though *v. Limbeck* and others deny this. According to *Winternitz*¹⁰² and *Thayer*,¹⁰³ a cold bath has the effect of increasing the proportion of leucocytes in the blood.

Leucocytosis can be readily diagnosed with a little practice by means of the microscope alone. For the more precise estimation of its degree the Thoma-Zeiss instrument may be used. It is necessary, above all things, that the blood examined should not be rich in digestion-products, and on this account it should not be taken until some hours after a meal. It is important to be able to recognise a pathological leucocytosis. Its presence taken in conjunction with the clinical symptoms,

will often serve to clear up the diagnosis of diseases—as, for instance, osteomyelitis—which, without its light, would be very obscure, and certain forms of pneumonia,¹⁰⁴ and will help to distinguish between typhoid pneumonia and septicæmia. From his observations in a number of cases—28 in all—the author concludes that in pneumonia, and to a variable degree in typhoid, a temporary and sometimes very marked leucocytosis may be produced by injecting pilocarpin, or by giving nuclein with the food.¹⁰⁵ *J. Schneyer*¹⁰⁶ found an increase of leucocytes constant in gastric ulcer and non-malignant pyloric obstruction, and regards the condition as diagnostic of the lesion.

3. Leukæmia.—This condition, when well marked, may be recognised from the naked eye characters of the blood (*Virchow*),¹⁰⁷ which,

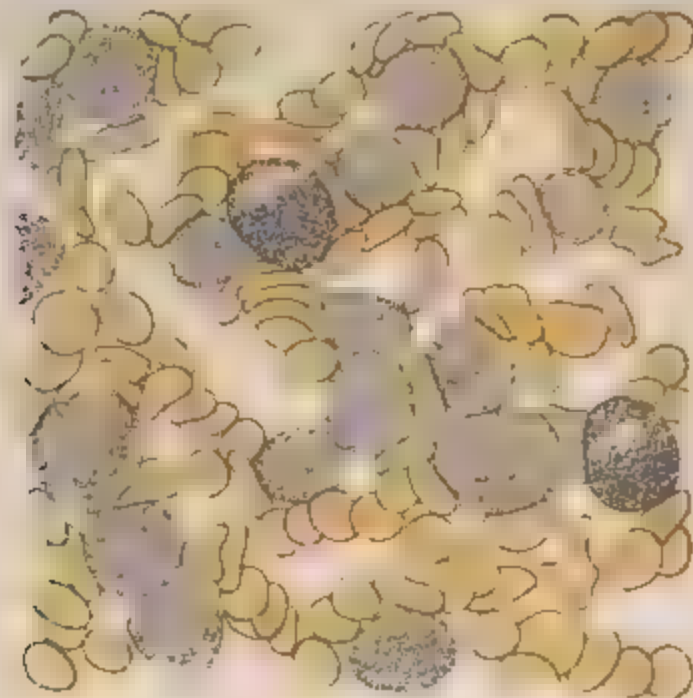


FIG. 16. Leukæmic Blood from a case of Splenic Leukæmia (shown by Zeiss compensation eye piece VIII ; objective, apochromatic immersion 1 : 30).

when allowed to flow from the finger, is light red in colour, and somewhat turbid and greasy, as though loaded with fatty matter.¹⁰⁸

The reaction of the blood is alkaline (*Mosler*),¹⁰⁹ not acid, as was formerly thought, but its alkalinity is often considerably less than normal (*v. Jaksch*). Microscopical examination shows an enormous increase in the number of white corpuscles [*J. H. Bennett*]. *Virchow* has found them in the proportion 2 : 3 of red corpuscles; *J. Vogl*, 1 : 3 to 1 : 2; *Schreiber*, 2 : 3.¹¹⁰ In five cases recently treated in Nothnagel's clinic, the proportion was 1 : 3, 1 : 5, 1 : 8, 1 : 11, 1 : 12 (*Gottlieb*).

In a typical case of leukæmia observed by the author in a child of sixteen months, the figures were, 1 : 40, 1 : 50, and 1 : 18;¹¹¹ and in nine cases recently observed of different forms of leukæmia the number

of leucocytes ranged between 163,000 and 992,000 to the cubic millimetre. The proportion of white to red corpuscles varied between 1 : 2.5 and 1 : 23.

Another notable characteristic is the diminution of the cellular elements of the blood in general. In the cases recorded above, their number fell to 2-3 millions in the cubic millimetre of blood, and in one instance to 1,510,000, the lowest point recorded in the observation of the nine cases referred to above. By the last examination in the child of sixteen months it was shown to be 2,440,000.¹¹¹

The amount of haemoglobin in the blood is usually diminished in leukæmia. In the case of the child mentioned above it was shown by v. Fleischl's haemometer to be 6.4 grms., and sank with the progress of the disease to 3.5 grms. In the series of nine cases it fell to between 10.5 and 1.12 grms.

It is important, further, to determine what is the prevalent character of the leucocytes present in the blood ; as to whether they are large or small, mono- or poly-nuclear. By observing this, the author believes, in spite of *Bizzozero's*¹¹² assertion to the contrary, that it is possible to ascertain with what form of the disease we have to deal. It is usual to distinguish leukæmia as splenic, lymphatic, and myelogenic, according to the anatomical seat and clinical symptoms of the disease ; although, probably, instances of true myelogenic leukæmia are very rare, and the anatomical distinction can seldom be strictly drawn, because the autopsy generally shows a greater or less degree of leukæmic change in all the organs examined.* [The existence of a myelogenic leukæmia is not recognised by *Hayem*.¹¹³ *Gowers*¹¹⁴ left the question open. Since he wrote, a case of undoubted myelogenic leukæmia has been described by *Dr. Wallace Beatty*¹¹⁵ of Dublin.]

When large and small leucocytes (lymphocytes)—the latter preponderating—are found in the blood, the leukæmia is of a lymphatico-splenic character. When, on the other hand, the larger cells alone are found, we are justified in concluding that the disease is splenic, with but little involvement of the lymphatic glands or marrow. If many corpuscles of a transitional form are found, nucleated red cells (normo-megalo-microblasts, *Ehrlich*), and especially large white multinuclear corpuscles holding eosinophil granules, and if, moreover, free eosinophil granules are in considerable quantity, there remains no doubt that the bone-marrow is the seat of serious changes, and the disease is of the myelogenic type.¹¹⁶

* Several years ago the author had under observation in Professor Nothnagel's clinic a case of nephritis in which the blood contained leucocytes of great size and with unusually large nuclei, their proportion to the red corpuscles (as determined by a single examination) being as 1 : 50. The autopsy revealed, in addition to chronic nephritis, changes in the medulla of the bones, like those described by Naumann in connection with leukæmia.

It must, however, be admitted that nucleated red cells may be found in cases of leukæmia which are not in any other way to be identified with the myelogenic disease (*v. Jaksch*), and in conditions which are not leukæmia at all (*Neumann*), whence it follows that these bodies are not pathognomonic. [Large and small leucocytes are found in every case of leukæmia. The smaller are most numerous in cases of moderate severity. When the disease tends rapidly to a fatal ending, very large hypertrophied leucocytes and even giant cells occur (*Hayem*. See also paragraph on anaemia infantum below.) Deviations from the normal in size are associated with a loss of contractility. The very large leucocytes, and also the smallest, fail to exhibit amæboid movements. Another peculiarity noticed by *Hayem* is the fact that the

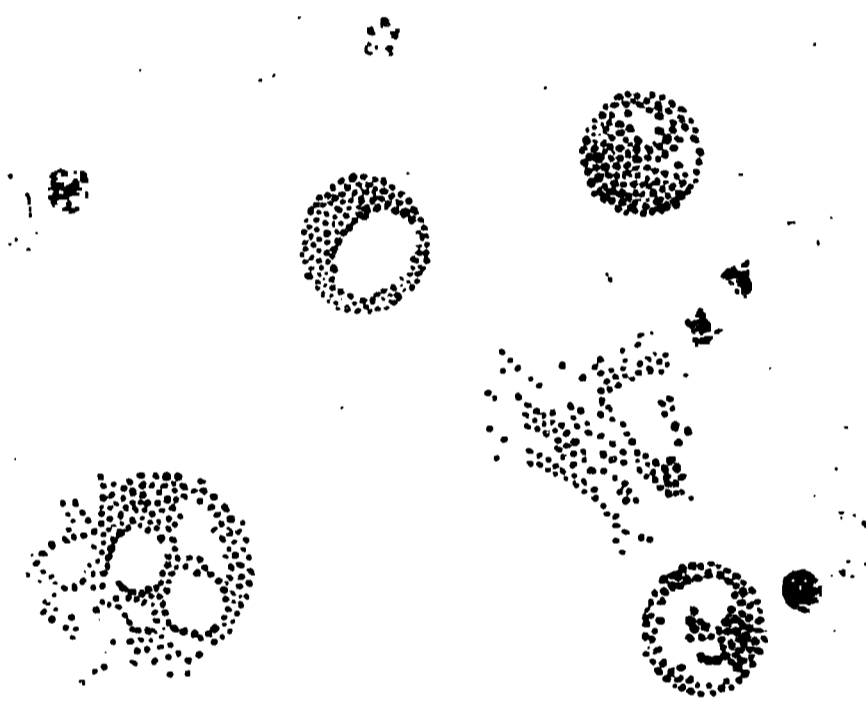


FIG. 17.—Eosinophil Cells from Leukæmic Blood (Zeiss's compensation eye-piece objective, apochromatic immersion 1.30; Abbe's reflector, open condenser).

leucocytes in this disease contain some haemoglobin more commonly than they do in health.^{117]}

Crystals are occasionally found in leukæmic blood (*Charcot, Robin, Vulpian*).¹¹⁸ *Neumann*¹¹⁹ attributes their origin to the marrow, and describes them as colourless, shining, oblong octahedra (*Ph. Schreiner*).¹²⁰ *Neumann*¹²¹ asserts that they are not present in the blood of splenic and lymphatic leukæmia. They are seldom met with. The author has often sought, but never seen, them in freshly drawn blood. It may be that they form only in blood that has been allowed to stand.¹²² *Prus*¹²³ has observed them in the freshly shed blood, and *Westphal*¹²⁴ in the living tissues of leukæmic patients.*

* For the chemical constitution of these crystals, see Chapters IV., VI., and IX.

It should also be mentioned that the red blood-corpuscles commonly undergo changes of form in leukæmia, a condition which first received the name of *Poikilocytosis* from *Quincke* (p. 41).

An advanced stage of leukæmia is readily recognised by the aid of the microscope, but it is by no means easy to distinguish between commencing leukæmia and a pronounced leucocytosis; and this is a task which the physician will often have to undertake. *Magnus Hues* and others¹²⁵ employ the term "leukæmia" only when the white corpuscles are in a proportion not less than 1 : 20. That the diagnosis of leukæmia cannot be made to depend upon this fact alone, however, is sufficiently proved by the author's investigation of cases of anaemia in children, where the proportion of leucocytes to red corpuscles was as 1 : 12, 1 : 17, and 1 : 20.¹²⁶ In another case, where the patient was an adult, the proportion stood at 1 : 7.3, and yet the condition was not one of leukæmia.¹²⁷ It is still more difficult to discriminate a temporary leucocytosis from the early stage of leukæmia.

We are indebted to *P. Ehrlich*¹²⁸ for an excellent means by which we may sometimes recognise an early stage of leukæmia. By observing the "granules" present in the protoplasm of the white corpuscles, he found that these uniformly exhibited remarkable differences in their staining properties—differences which have both a physiological and a pathological significance. Upon this basis he distinguishes five several varieties of what he calls "granules," classing them as α to ϵ granules. In all cases of acute leucocytosis, the mono- and poly-nuclear forms which furnish the ϵ granules are alone increased in number, whilst the α granules—which are also styled "eosinophil," from their property of taking up eosin—are apparently fewer. Precisely the reverse of this, again, obtains at the beginning of leukæmia: the eosinophil granules are increased in quantity, as also, according to *Ehrlich*, are the basophil cells. The method of examination is as follows:—The blood is spread in a very thin layer between two cover-glasses, which are then grasped with forceps, dried in a exsiccator, and heated upon plates of copper foil (for this purpose a drying chamber for temperatures beyond 100° C. will serve very well*) for a considerable time (10 to 12 hours) at 120°-130° C. A drop of concentrated eosin-glycerine solution is then added to the preparation, the colouring matter is washed off with water, and the preparation dried, and examined in Canada balsam or oil of cloves.

According to *Huber*,¹²⁹ good results may also be obtained in the following manner:—Two grms. each of aurantia, indulin, and eosin are dissolved in 30 grms. of glycerine, the resulting thick fluid well shaken

* Hardening with absolute alcohol may also be employed; then, of course, it will be necessary to drive off the alcohol before staining.

up, and the cover-glasses (dried and heated as above for several hours at 120° C.) are immersed in it for a period varying from half-an-hour to some days. When taken out, they are carefully washed with distilled water, dried in the air, and examined in Canada balsam or dammar varnish. Should the case be one of early leukæmia, the red blood-corpuscles in such a preparation are stained a reddish-yellow, the nuclei of the white corpuscles have taken up the blue colouring matter, and in addition there are seen large leucocytes (eosinophil cells) distended with granules of a deep red tint (eosinophil granules, fig. 17). Sometimes the whole field is covered with these granules. In such preparations are also commonly to be seen large colourless cellular bodies of oval shape, distended at their poles with similar particles.

*Gabritschewsky*¹³⁰ and *Aldehoff*¹³¹ stained blood-preparations made as above with eosin. The latter employed a concentrated alcohol (bluish) solution of eosin, that known as No. 22 of Bayer's factory at Elberfeld, in the following way :—The preparation is submitted to the action of the staining fluid for half-an-hour in the cold, or for two to three minutes when heat is used ; the excess of colouring matter is then washed away with distilled water, and the preparation is placed for a short time in a concentrated watery solution of methylene-blue, dried and examined in Canada balsam. Very beautiful specimens may be obtained in this way. Sometimes the preparation will display the process of karyokinesis in the red corpuscles (fig. 17).

Important modifications of these methods have been lately introduced. The following reagents have been employed :¹³²—

Ehrlich's Triacid Mixture.—This consists of 115 cc. of a saturated watery solution of orange-green ; 125 cc. of a saturated watery solution of acid fuchsin to which 20 per cent. alcohol has been added. To this are added 125 cc. of a saturated watery solution of methyl-green and 77 cc. absolute alcohol, the mixture being well shaken the while.

Neusser's Reagent¹³³ is :—

Saturated watery solution of acid fuchsin	50 cc.
Orange-green	70 cc.
Methyl-green	80 cc.
Absolute alcohol	80 cc.
Glycerine	20 cc.
Distilled water	150 cc.

A series of observations which the author has made with the blood of healthy and anaemic persons, and especially of rickety children, has shown that the appearances in question occur but rarely in normal blood or in that of anaemic states. In one case only, that of a patient with tuberculosis, and who was not leukæmic, were they to be seen in abundance. *Aldehoff* observed a large number of eosinophil cells in

the blood in three cases of malaria; so also did *Dolega*.¹³⁴ The author has found such in healthy adults in pneumonia, and in anaemia of all kinds. *Müller* and *Rieder*¹³⁵ have had the same experience. *Fink*¹³⁶ has described numerous leucocytes with eosinophil granules as present in the blood of asthmatic patients.

It follows that the detection of eosinophil granules in increased abundance has lost much of its weight as an evidence of commencing leukæmia. Still the increasing observations of *Müller*¹³⁷ afford room for the belief that, by more accurate differentiation of eosinophil leucocytes, further conclusions may be reached; and he has shown that in leukæmia there is probably but one form (Cornil's "mark-zellen") to be found which does not occur also in healthy blood. The matter will have acquired a diagnostic importance only when it has been proved conclusively that the bodies in question do exist in the blood of leukæmia alone, and do not exist in that of other forms of anaemia. For the present the assumption is not warranted. The author has observed leucocytes similar to those described by Müller in the blood in a case of sarcoma, and *Weiss*¹³⁸ has reported in like manner. Nevertheless, where repeated observation has shown the presence of these bodies, it is perhaps safe to admit them as evidence of commencing leukæmia.

The observations of *Neusser*,¹³⁹ *Zappert*,¹⁴⁰ and others of late years have drawn much attention to the subject of the eosinophil cells, and a great clinical interest has attached to it. Further experience is needed to decide how far the very suggestive conclusions drawn by *Neusser*¹⁴¹ correspond with the facts. The same writer's "perinuclear basophilia" is still under discussion.

In view of the great importance of *Ehrlich's*¹⁴² process for the investigation of the blood, some further notice of his methods is called for. [1. *Eosinophil* granules are those which stain with acid pigments, of which eosin is one. The process for their detection has been already described. 2. *Basophil* granules stain with basic aniline dyes, e.g. dahlia, gentian-violet, methyl-violet, methyl-green, vesuvin, and fuchsin. 3. *Neutrophil* granules stain best with neutral dyes, i.e. those composed of a coloured base and an acid. Of these, methyl-blue and acid fuchsin are examples.¹⁴³]

[S. *Dépine* divides the aniline dyes into two classes. (1.) Acid fuchsin may be taken as a type of the first; (2.) basic fuchsin as a type of the second. The first class stains deeply the most differentiated parts of the cells; the second, the least differentiated parts of the cells. In a drop of blood, the red corpuscles are most deeply stained by acid fuchsin, the leucocytes but faintly stained, while the nuclei are not stained at all. If basic fuchsin be used, the nuclei are stained most deeply, the body of the leucocytes to a less extent, and the red corpuscles least of all.¹⁴⁴]

For the detection of *neutrophil* or ϵ -granules, a fluid is required of the following composition:—To five parts of a saturated solution of acid fuchsin one part of a watery solution of methyl-blue, and five parts of distilled water are added; the mixture is allowed to stand for some days, then filtered, and the filtrate may be employed for staining blood-preparations in the manner indicated above. The leucocytes will then exhibit deep violet-coloured granules.

For the detection of basophil or γ -granules ("mast-zellen" granules*) a fluid is needed containing 50 cc. of a saturated alcoholic solution of dahlia and 10 cc. of glacial acetic acid in 100 of water.¹⁴⁵

According to Ehrlich, basophil granules are absent from the blood in health. It is probable that in the near future leucocytes may be further distinguished according to their development, and the information so obtained may become available for the purposes of diagnosis, as Einhorn,¹⁴⁶ Zappert,¹⁴⁷ Elzholz,¹⁴⁸ and Thayer¹⁴⁹ suggest. It would be of great interest to ascertain which of the recognised varieties of leucocytes occur in the blood in the conditions of leucocytosis mentioned above. According to Einhorn, they would appear to be especially the polynuclear forms.

It remains to mention that leukæmic blood commonly contains peptone. (See p. 82.)

4. Anæmia Infantum Pseudo-leukæmica.¹⁵⁰—The author has

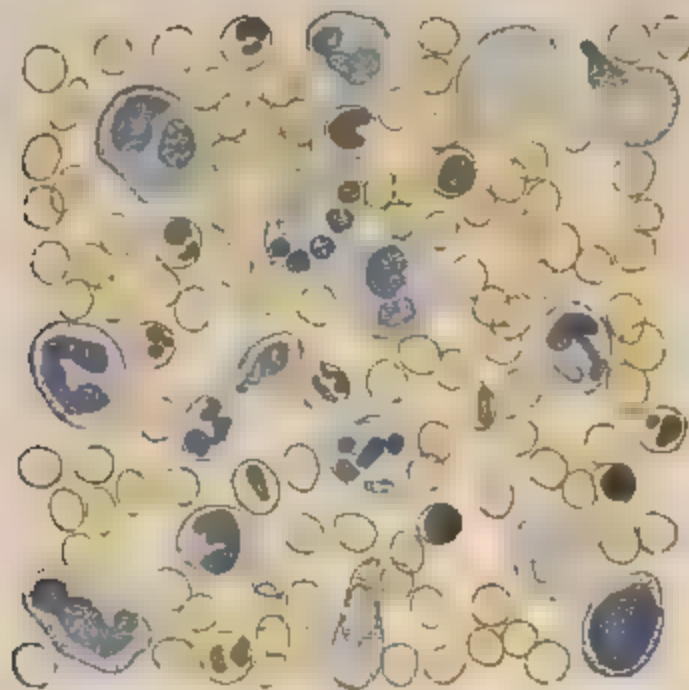


FIG. 18. Appearance of the Blood from a Case of Anæmia Infantum Pseudo-leukæmica. (Zeiss's compensation eye-piece 4, achromatic objective $\frac{1}{2}$, homogeneous immersion, Abbe's mirror and open condenser. The blood was stained by Becker and Huber's method.)

described a form of anæmia of a very distinctive character as occurring in children, and the observations of Loos¹⁵¹ in connection with the blood changes concerned, and of Luzet,¹⁵² Hock and Schlesinger,¹⁵³ Monti and Berggrun¹⁵⁴ in respect of its clinical history, furnish sufficient warrant for regarding it as a separate disorder. Pathologically the most striking fact is a very great diminution of the colouring elements of the blood. In one case the red corpuscles numbered only 820,000,

* ("Mast-zellen" is the name given by German authors to certain bodies which occur chiefly in connective tissue in the neighbourhood of blood-vessels. They are round, nucleated, larger than leucocytes, and they have granules embedded in their protoplasm. These granules stain deeply, and have been mistaken for micrococci.)

the white 54,666. Leucocytes are always increased in number, but the increase is not so great or so progressive as in leukæmia. Further, the leucocytes display a remarkable variety of form and attain to an unusual size. The red corpuscles in freshly shed blood exhibit a high degree of poikilocytosis (see p. 41), and their contents are apt to be colourless—a condition which is perhaps a form of poikilocytosis (compare figs. 20 and 21).

[*Hayem*¹⁵⁵ has already commented upon the appearance in leukæmic blood of red corpuscles of pale colour and devoid of hæmoglobin. These, he says, are not merely stroma, and he inclines to the belief that they are stroma *plus* an albuminoid material with which hæmoglobin normally combines.]

Some of the leucocytes hold red corpuscles or fragments of the same embedded in their substance, and amongst them are a few basophil, and very large polynuclear neutrophil cells. Finally, nucleated red corpuscles are also to be met with.

It is to be observed that none of the characters mentioned here as belonging to the blood in anæmia infantum pseudo-leukæmica are peculiar to that condition. Nucleated red corpuscles, for instance, are found in leukæmia, pernicious anæmia, and purpura (*Spietschka*¹⁵⁶). The author lately observed in a case of rheumatic pericarditis with profuse hæmorrhage (*Peliosis rheumatica*) an unusually large number of nucleated red corpuscles; indeed, every preparation of this blood had a large proportion of them.¹⁵⁷

It is well known that the red corpuscles are all nucleated during early foetal life, and, according to *Hayem*,¹⁵⁸ begin to be replaced by the coloured non-nucleated erythrocytes or red cells only in the seventh month.

There are, however, clinical symptoms—enlarged spleen, &c.—a description of which would be out of place here, and these, together with the character of the blood, sufficiently distinguish the disease.

It may be mentioned that the changes in the blood are very similar to those found in leukæmia, but in the latter the cellular elements and the amount of hæmoglobin are never reduced so much.

That the condition in question occurs in childhood and is evidenced by definite symptoms, is attested by various observers.¹⁵⁹

5. Melanæmia.¹⁶⁰—The microscopical appearance of the blood in this very rare disease suffices for its recognition. There are to be seen floating amongst the blood-corpuscles smaller or larger granules and granulation masses, possibly pigment lumps, which are usually black, more seldom brown and yellow. The granules are united to each other by a substance which is soluble in acids and alkalies. The pigment occurs also in the form of separate granules equal in size to leucocytes.

Finally (and this, in the author's opinion, is the commonest case), the pigment particles, both small and large, may be enclosed within cells which occasionally resemble white blood-corpuses, and sometimes differ from them in being flask- or spindle-shaped. The pigment lumps are very rarely observed ; but, on the other hand, the granules, and, more commonly still, pigment-laden white corpuscles, are often temporarily present after a severe attack of ague and in relapsing fever. The preparation (shown in fig. 19) was made from the blood of a man who had suffered for a year from malarial disease which he had contracted in the tropics. The appearances described here are usually associated with oligochromæmia and oligocythæmia, and accompanied by the symptoms of anaemia (compare p. 44).

6. Microcythæmia.—This condition was first described by *Vanlair* and *Masius*.¹⁶¹ It is characterised by the presence in the blood of small haemoglobin-containing elements (*microcytes*), which are probably derived

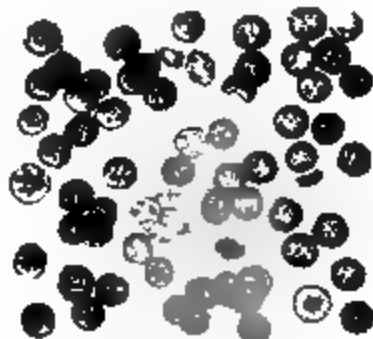


FIG. 19.—Melanemic Blood (eye-piece III., objective 8A, *Reichert*), from a Case of Malarial Cachexia.

from the red blood-cells, and are generally smaller, but occasionally larger (*megaloblasts* of *Hayem* and *Ehrlich*), than these.

Such bodies are found in many morbid states, and notably in toxic states, infectious diseases, pronounced anaemia, and in burns. Notwithstanding that much has been written about them, there is really but little known on the subject of microcytes, and their discovery is of no assistance in diagnosis. *Litten* has shown that they may rapidly appear in the blood, and as rapidly disappear from it. We may refer here also to *Bettelheim's*¹⁶² observations on the subject of minute mobile granules in the blood. *Gram* and *Grüber*¹⁶³ regard the microcytes as the result of post-mortem changes in the blood. The latter believes that they represent the final product of the rapid and uniform abstraction of water from the corpuscles ; and since such a process will take place most readily in blood which contains but little water (and is, therefore, relatively rich in albumin), the subject of microcythæmia is invested, by this theory, with a certain practical importance.

7. Poikilocytosis.—By this term it is usual to describe an affection of the blood in which the red corpuscles exhibit a remarkable *variety* as to form and size.¹⁶⁴ The condition was first noticed in pernicious anaemia, and has been regarded by some authors as pathognomonic of that disease. Cases, however, have been recorded (*Grainger Stewart, Lépine, Hermann Müller*) in which no such changes were found.



FIG. 20.—Poikilocytosis from a Case of Amyloid Degeneration of the Kidney, Liver, Spleen, and Intestine (eye-piece III., objective 8A, *Reichert*).

The appearance of the red blood-corpuscles in this condition exhibits a remarkable variety. Some are normal in shape and size, but amongst them are seen smaller forms and others of great size (megaloblasts). Some, again, are flask-shaped, and often tipped at one extremity with a little knob; while others of irregular outline have been compared to such different objects as an anvil, a biscuit, a goblet, or a kidney

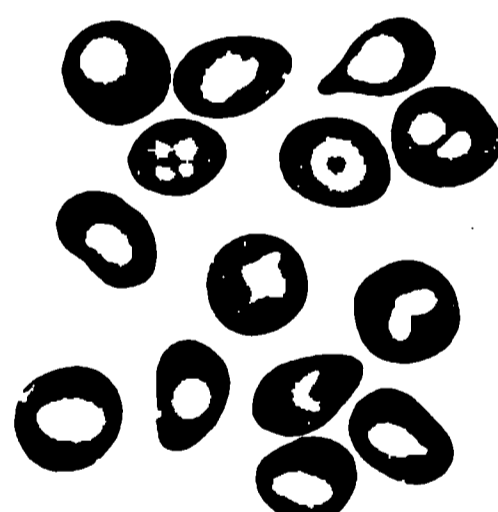


FIG. 21.—Blood in Poikilocytosis (compensation eye-piece 4, *Zeiss's apochromatic objective*), from a Case of Anæmia Infantum Pseudo-leukæmica.

(fig. 20). *Friedreich* and *Mosler* have seen amoeba-like processes on the red corpuscles. The author has also observed similar appearances, and inclines to the belief that the prominent features of poikilocytosis depend upon the fact that the red corpuscles are endowed with an abnormal power of contractility. This quality may also belong to the contents of the red corpuscles¹⁶⁵ (see fig. 21), and it is possible that the contained substances lately noted within them in cases of cancer may be

accounted for in this way.¹⁶⁶ *Quincke*¹⁶⁷ has observed this condition, and he has applied to the resulting appearances the name of "cup-shaped depressions in the red blood-corpuscles." From the above description it may be seen that poikilocytosis is easily distinguished with the microscope. Such appearances are not confined to any one disease, but are common to many in which the blood is greatly altered, and especially in respect of the diminution of its cellular elements and of the red corpuscles in particular.

The author has himself observed them in cases of chlorosis and severe anaemia of whatever kind, and typically developed in pernicious anaemia, and in anaemia infantum pseudo-leukæmica. He has also seen them in the cancerous cachexia, in leukæmia, in amyloid degeneration of the viscera, and in cases of chronic nephritis. *Maragliano*¹⁶⁸ maintains that poikilocytosis is in reality a necrosis of the red corpuscles, and that it indicates a formidable and always eventually fatal disorder. The author's own observations enable him to concur in *Maragliano's* view, but with this limitation, that the condition may be present in chlorosis without involving such serious consequences. *Gräber*¹⁶⁹ believes that the poikilocytes do not exist in the circulating blood; a view which the author does not assent to in all cases.

8. Changes in the Formed Elements of the Blood in Chlorosis.¹⁷⁰—Although the minute examination of the blood in chlorosis commonly discloses few or none of the changes previously described, it will prove both interesting and profitable to mark those particulars in which it differs from simple oligocythaemia on the one hand, and from pernicious anaemia on the other.

Chlorotic blood is especially characterised by its paler colour, not necessarily accompanied by other changes in the physical properties of its constituents. Microscopical examination shows usually that the red corpuscles are abnormally pale without their number being markedly diminished. The application of the methods for counting the corpuscles and for estimating the amount of contained haemoglobin proves, in the greater proportion of cases, that while the number of corpuscles is but slightly altered, the blood is very deficient in haemoglobin.¹⁷¹ In a series of eight cases of typical chlorosis investigated at the beginning of treatment, the red corpuscles ranged between 6,300,000 and 2,684,000, and leucocytes between 14,000 and 6600 to the cubic millimetre of blood, while the haemoglobin was reduced proportionally much more. In one instance it reached 7.8 grms. in 100 grms., but was generally between 5.6 and 2.8 per cent. Generalising somewhat, the conditions that obtain in chlorosis may be said to be:—Diminution of the amount of haemoglobin, together with a less diminution of the cellular elements of the blood, and with or without an accompanying relative increase

of leucocytes. Poikilocytosis is a frequent phenomenon in chlorotic blood, whilst microcytes, megaloblasts, and nucleated red corpuscles are also not of rare occurrence. In some cases of chlorosis the number of red corpuscles is so greatly diminished that they approach the character of pernicious anaemia in this particular. The remarkable case recorded by *Luzet*¹⁷² may have been of this order. Finally, according to Peiper and Gräber, the alkalinity of the blood is always increased. The author has noted a diminished alkalinity in three instances.

9. Changes in the Formed Elements of the Blood in Pernicious Anaemia.¹⁷³—Very different is the condition witnessed in pernicious anaemia. The blood is thin, light-coloured, and shows all the changes which have already been shown to belong to oligocythaemia (p. 8). Under the microscope its cellular elements are seen to be enormously diminished in quantity, a condition which is seldom or never observed in the very worst forms of simple anaemia. The number of blood-cells, according to *Laache*, may fall so low as 360,000 to the cubic millimetre. *Sadler*,¹⁷⁴ working in the author's clinic, obtained 872,000–562,000 red corpuscles as the lowest numbers; and in three cases of pernicious anaemia verified by autopsy the numbers were 971,875, 512,000, and 512,000. The enumeration was made in these cases a few days or hours before death, and they go to prove that a fall to below a million in the number of the red corpuscles is a fact of gloomy significance. Further, the individual red cells are often larger than normal, and exhibit in an exquisite manner the characteristics of poikilocytosis. Eosinophil cells are also present in unusual number, and the characters belonging to severe anaemias generally and to anaemia infantum pseudo-leukæmica may be observed, except that the degree of leucocytosis is never permanently so great as in the latter disease. Nucleated red corpuscles, *Ehrlich's* megaloblasts, and karyokinetic arrangement in the contents both of white and of red corpuscles are not wanting.¹⁷⁵

To estimate the diameter of the corpuscles, the method of "dry measurement" may be employed. It is founded upon the fact observed by *C. Schmidt*, that red blood-corpuscles, when rapidly dried, retain their form permanently. According to *Laache's* directions, the proceeding is as follows:—A slide is gently heated and then swept quickly and lightly over the surface of a very small drop of blood drawn from the skin. The blood dries rapidly, and when placed under the microscope, the corpuscles appear as bi-concave discs, lying apart from one another, so that their diameter can easily be measured with the familiar micrometric apparatus (micrometer eye-piece). In the case of healthy blood corpuscles, the diameter ranges between 6.5 or 6.7 and 9.0 or 9.4 μ .*

* The Greek letter μ represents one-thousandth of a millimetre ($\mu = 0.001$ millimetre), and is the sign of a micromillimetre or micron.

In the blood of pernicious anaemia, red corpuscles are found of $10-15 \mu$ diameter. Microcytes are seldom found in the blood in this disease.

An important sign of pernicious anaemia, as *Hayem* first showed, is the fact that the number of red cells in the blood is inversely proportional to the quantity of haemoglobin which they contain. *Copeman*¹⁷⁶ made the remarkable discovery that when the blood of pernicious anaemia is quickly dried, rhombic crystals of haemoglobin are sometimes formed.

The presence of amorphous haematoxin in fresh blood is not a very rare occurrence. The author met with it in the case of a child of four months suffering from congenital syphilis and severe icterus.

The author has shown that with the progress of the disease the blood of pernicious anaemia becomes richer in nitrogen.¹⁷⁷ Since the statement on this head was made, a series of observations on a case of the disease, subsequently verified by the autopsy, has confirmed the existence of a hyperalbuminaemia rubra. The result of these observations is given in the following table. In each case the figure taken is the mean of two estimations.

The patient was a lithographer, aged 36.

Date.	100 grms. Moist Red Corpuscles yielded.	Corpuscles=per cubic millimetre.		Hæmoglobin.
		Red.	White.	
Apr. 26, 1894	5.74 grms. N = 35.79 grms. albumin	980,000	7,600	3.5
May 5 "	5.91 " " = 36.95 " "	612,000	5,800	2.8
May 10 "	7.32 " " = 45.30 " "	512,000	6,000	1.75

In a healthy man the moist red corpuscles yield 5.52 per cent. nitrogen = 34.5 per cent. albumin.

We see then that the characteristic features of the blood in pernicious anaemia are : a general diminution of its cellular constituents, with a relative increase in the size of the red corpuscles and in the quantity of haemoglobin and of nitrogen which they contain.¹⁷⁸ All these properties can be shown by the methods described above and on p. 77.

[It is often important to have the means of comparing specimens of blood, whether from different individuals or from the same individual at different times. *Dr. R. Muir*¹⁷⁹ has adopted the following method for making permanent preparations of blood. A film of blood is obtained in the usual way upon a cover-glass, and while it is still wet the latter is placed in a saturated watery solution of corrosive sublimate, to which a little salt has been added, and is left there for thirty minutes. It is then removed, and the corrosive sublimate is washed off with a $\frac{1}{2}$ per cent. solution of common salt. The cover-glass is placed for a few minutes in spirit, and afterwards in absolute alcohol. The specimen can then be stained with eosin and logwood, and is absolutely permanent.]

Micro-organisms have been demonstrated, both by staining and by cultivation, in the blood of pernicious anaemia (*Fr. Fischel, R. Adler*¹⁸⁰). Of three typical cases verified by autopsy, the author found cocci in one

only ; and in this there may have been an error, as his colleague, *H. Chiari*, examined the blood post-mortem without result. It may be that different conditions are described by the name of pernicious anaemia, and that among them is a crypto-genetic septicæmia (*Fischel, Adler*), which bears a close resemblance to it.

10. Changes in the Formed Elements of the Blood after Hæmorrhage and in Infectious Diseases (Secondary Anaemia).

—In these conditions, and also in chronic nephritis, anaemia, with a diminution in the corpuscular elements and of the colouring matter of the blood, is of constant occurrence, but the other changes which have been shown to accompany pernicious anaemia, &c., are absent. *Neubert*¹⁸¹ has shown that in pulmonary phthisis the amount of haemoglobin frequently diminishes more rapidly than the number of corpuscles.

In myxœdema, *Kræpelin*¹⁸² observed blood-changes similar to those which belong to pernicious anaemia, and the same microscopical appearances have been found in cases of anaemia due to the presence of intestinal worms, as *Bothriocephalus latus*, *Dochmias duodenalis* (see Chap. VI.), and in the course of syphilis.¹⁸³ In congenital syphilis, according to *Loos*,¹⁸⁴ the anaemia is so severe as to be sometimes the immediate cause of death. The results of malarial infection have been already noticed (p. 40).

V. THE PARASITES OF THE BLOOD.—Amongst these are members both of the vegetable and of the animal kingdom.

A. Vegetable Parasites.—We shall follow the plan usually adopted in clinical medicine, by which micro-organisms are divided into three great groups—(1) Moulds; (2) Yeasts (*Saccharomycetes*); and (3) Fission-fungi (*Schizomycetes*, bacteria). The third group alone concerns us here, for to it belong, almost without exception, the parasites which as yet have been found to infest the blood. Yeasts are mentioned in an observation of *Busse's*,¹⁸⁵ and moulds have occasionally been seen in the blood of animals,¹⁸⁶ but not yet in that of man, nor has any definite disease been attributed to their agency. We have, therefore, to describe the bacilli of *anthrax*, of *relapsing fever*, of *tubercle*, of *glanders*, and of *typhoid*, and also the cocci, streptococci, and staphylococci which infest the blood. With reference to the bacillus of *tetanus*, the recent investigations as to its occurrence and significance are to be received with great reserve. In view of the researches of *Kitasato*,¹⁸⁷ however, it is no longer possible to question the pathological significance of this micro-organism. The statements concerning the presence of streptococci in the blood of certain diseases also need confirmation before they are made the basis of clinical inferences.

Methods of Examining the Blood for Micro-Organisms.—In certain

diseases (as, e.g., relapsing fever and anthrax) no difficulty will be found in determining the presence of bacilli in the blood by means of a simple microscopical examination. In other instances (and especially in miliary tuberculosis, typhoid, and glanders) we must resort to the methods of *Koch* and *Ehrlich*. In all alike the common process is to dry the blood in a thin layer, so that the shape of the corpuscles is lost without altering the appearance of the bacilli, and then to submit it to the staining methods employed by *Koch*, *Ehrlich*, *Weigert*, &c.,¹⁸⁸ and other investigators. *The general principle of the methods is this, that the fungi are deeply stained by the basic aniline dyes.* To these basic aniline dyes belong: Bismarck-brown, vesuvin, aniline-brown, fuchsin, methylene-blue, gentian-violet, and methyl-violet. It must be observed, however, that other substances besides the micro-organisms are coloured by these dyes. Thus, for instance, masses of protoplasm, cell-nuclei, and the products of cellular disintegration stain readily with them. This is true also of the γ and δ granules of *Ehrlich*, and, in fact, they have over and over again been mistaken for fungi.

Preparation of Cover-Glasses.—The skin of the finger-tip from which the blood is to be taken must first be thoroughly cleansed with soap and water and a nail-brush, and washed with corrosive sublimate (1 in 1000). All traces of the latter are removed by first rinsing the finger with alcohol, and finally pouring æther upon it. A pretty deep puncture is now made in it with a needle previously sterilised by heat. The small instrument devised by *Hauxley*¹⁸⁹ may be used instead of the needle.

Scheurlen's process is, perhaps, safer, although, as a fragment of glass may remain in the wound, it is not free from danger. Numerous observations have shown that aseptic blood can be obtained in the ways mentioned here. *R. Kraus* strongly advocates the obtaining of the blood from a vein.¹⁹⁰

The first drops of blood that issue are swept away with a well-sterilised platinum needle, and a cover-glass held in steel forceps, also sterilised, is quickly brought in contact with the surface of the blood as it flows freshly from the wound. To this cover-glass another is at once applied, and the blood spread in a very thin layer between them. They are then separated with the aid of two pairs of forceps, and placed to dry in an exsiccator or in a chamber free from draughts and dust. Care should be taken that the cover-glasses employed are absolutely free from impurities, and to this end they should have been washed before use in corrosive sublimate, alcohol, and æther. Not less care must be taken to secure the cleanliness of the needles and forceps used. When dried in this manner, the cover-glass, with the prepared surface upwards, is three times passed cautiously through the flame of a Bunsen's burner, and

then kept for some hours at a temperature of 120° . After this a drop of a strong watery solution of a basic aniline dye is applied to it by means of a pipette, allowed to rest on it for a short time—some minutes at the longest—and then washed off with a gentle stream of sterilised distilled water, directed so that it flows over, but not directly down upon, the stained parts. With this object the cover-glass should be held obliquely to the stream. The preparation can be examined whilst wet. To mount it in Canada balsam, dammar varnish, or oil of cloves, it is necessary to dry it again before placing it with a drop of one of these substances on a slide.

If too concentrated a staining fluid has been used, and the specimen proves too highly stained, the excess can be removed by the addition of alcohol.* *Ehrlich*¹⁹¹ claims for methylene-blue the advantage that, even when submitted to its action for a long time, the preparation will not be too deeply coloured.

Over-staining is less likely to occur when alcohol, glycerine, or acetic acid has been added to the water in which the dye is dissolved. It must be mentioned, however, that vesuvin, Bismarck-brown, and aniline-brown cannot be employed in alcoholic solution.

Further, it is a good plan, especially for the purposes of a provisional inspection, to treat the cover-glass directly by placing upon it a drop of a filtered alcoholic solution of an aniline dye, such as fuchsin or methyl-violet, and then to wash off the superfluous staining fluid with alcohol, when the preparation may be examined in the manner already described.

With reference to the staining fluid, it is advisable to prepare it afresh each time it is required, because it readily decomposes when allowed to stand; and besides, fungus vegetation is apt to thrive rapidly in dilute solutions. To examine a dry cover-glass preparation of blood for micro-organisms, the directions of *Löffler*¹⁹² may be followed with advantage:—The cover-glasses, prepared as above, are left for 5–10 minutes in a staining fluid consisting of 30 cc. of a concentrated alcoholic solution of methylene-blue, and 100 cc. of 1 : 10,000 solution of potash, then washed for 5–10 seconds in $\frac{1}{2}$ per cent. solution of acetic acid, treated with alcohol, dried, and mounted in oil of cloves or Canada balsam.

*Gram's*¹⁹³ method, which is also to be commended for the examination of micro-organisms, is somewhat different. He prepares the cover-glass in the manner described above, and places it for some minutes in an *Ehrlich-Weigert* solution of gentian-violet and aniline water (comp. Chap. IV.). When sufficiently stained, he puts it into a solution of iodine and iodide of potassium (iodine 1.0, iodide of potass. 2.0, distilled water

* Glycerine or dilute acetic acid will serve this purpose also.

300.0), when a dull red-brown precipitate forms. It remains in the iodo-potassic iodide solution for two or three minutes. It is then put to bleach in absolute alcohol, where it remains for some time. When taken out and examined, all the cellular elements are seen to be colourless, while the micro-organisms are deeply stained a dark blue.

In Weigert's¹²⁴ modification of the same process the cover-glasses are stained in a saturated aniline water gentian (or methyl-) violet solution, washed with water or normal saline, and dried. A drop of Lugol's fluid is then applied. When this has taken effect, the preparation is once more dried, and aniline-oil added, a drop at a time. The latter is then thoroughly washed off with xylol and the preparation examined in the usual way. By this method micro organisms are stained a dark blue, while fibrin particles have a pale colour.



FIG. 22.—Anthrax Bacilli from a Rabbit's Blood, from a preparation by Professor Weichselbaum
eye-piece III., objective Reichert A., homogeneous immersion, Abbe's mirror and open condenser).

Gunther's method of investigation for the spirillum of relapsing fever (see p. 51) is also applicable to the examination of blood for micro-organisms. The microscope employed in these researches should be provided with an oil-immersion lens, and an Abbe's reflector and open condenser (Chap. X.). The apochromatic objectives of *Reichert*, *Zeiss*, and other instrument-makers serve still better (see Chap. X.), while the semi apochromatics of *Reichert* are valuable for daily use at the bedside.

1. **Bacillus of Anthrax**—The presence of micro-organisms in the blood of men and animals suffering from anthrax was pointed out by *Pollender*, *Brauell*, and *Davaine*. Many other observers (*Buhl*, *Waldeyer*, *E. Wagner*, and *W. Müller*¹²⁵) have since described the occurrence of the bacillus of this disease in human blood; but in all cases the number as seen in man falls far short of that witnessed in the case of the lower animals, and it varies according to the part from which the blood is taken. It is found most abundantly in splenic blood.

When looked at through the microscope, the micro-organisms are seen to be motionless, rod like bodies, $5\text{--}12\ \mu$ long,* and almost uniformly $1\ \mu$ broad, slightly thickened at the extremities, and occasionally having the appearance of transverse segmentation towards the middle. Even in unstained preparations they can be readily seen when they are present in great numbers.

The blood is at the same time thin and of a dark red colour; commonly, too, pronounced leucocytosis coexists. Where typical anthrax bacilli are found in the blood, the diagnosis of the disease is established beyond question; but, on the other hand, it must be borne in mind that the symptoms of anthrax may be well marked while no

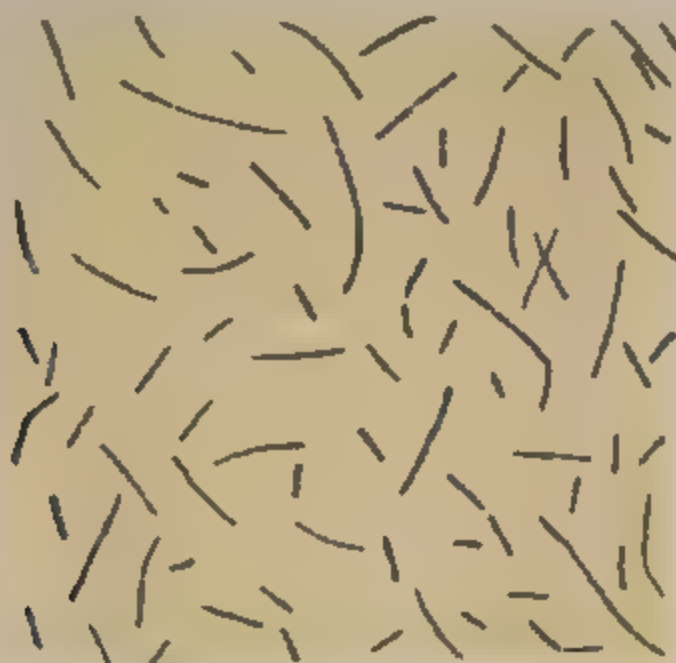


FIG. 23. Anthrax Bacilli from Human Blood (*post-mortem*), from a preparation by Professor Eppinger (eye-piece III, objective Zeiss $\frac{1}{2}$, homogeneous immersion; Abbe's mirror and open condenser).

bacilli are anywhere to be seen. In such a case the diagnosis may be established by injecting some of the suspected blood into an animal (mouse or guinea-pig), which, if anthrax be present, will shortly exhibit symptoms of the disease, and its blood when examined will be found to hold bacilli in profusion (fig. 22). The bacillus of anthrax does not form into long threads either in the blood or in the living tissues, and it has no spores (*R. Koch*).¹⁹⁸ It increases only by division. Fig. 22 represents bacilli from a rabbit's blood.

Fig. 23 is from a preparation made from the blood of a man who had died of anthrax (wool sorter's disease). A comparison of figs. 22 and 23 will enable the reader to recognise the bacilli.

The examination of the blood in anthrax is conducted in the manner

* $1\ \mu = 0.001$ mm.

described above (for dry cover-glass preparations and staining with basic aniline dyes). Löftler's method is especially applicable to the purpose. It should be mentioned here that the affection known as "wool-sorter's disease," the nature of which was formerly so obscure, is now known (from the observations made by *Eppinger* and *R. Paltauf*¹⁴⁷) to be identical with anthrax, and when the clinical symptoms of that disease come under observation, the physician should in all cases examine the blood and pathological fluids (pleuritic exudation, &c.) for bacilli.

[*S. Martin*¹⁴⁸ has distinguished a highly toxic alkaloid in the products of *Bacillus anthracis*, and has identified this in the blood and organs of wool-sorter's disease. He believes that this substance results from decomposition in the spleen and elsewhere of more complex albumoses. The alkaloid is more noxious than the albumose from which it is derived.]

2. Spirillum of Relapsing Fever.—The spirillum was first noticed by *Obermeye*¹⁴⁹ in the blood of a patient suffering from relapsing fever.

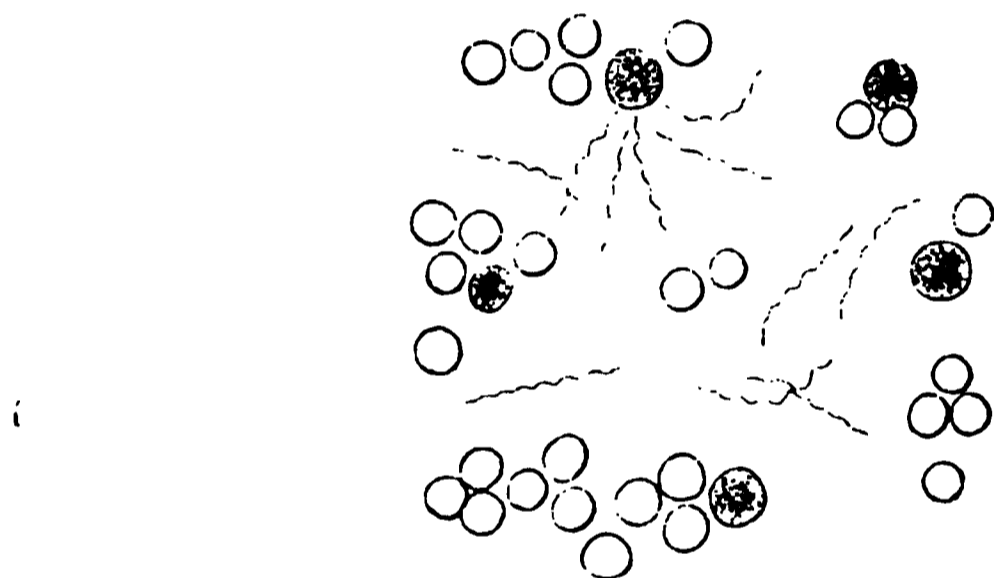


FIG. 24.—Spirilla of Relapsing Fever, from a photograph by Koch.

It has since been seen by many observers, but most authorities are agreed that it is to be found only during the paroxysms of the disease, and that as the temperature falls the spirilla disappear.* When a specimen of blood containing them is placed under the microscope, the spirilla appear as long and very delicate unsegmented threads twisted into spirals. Their average length is about six or seven times the diameter of a red blood-corpuscle. They have a brisk vibratile movement in the direction of their long axis. This motion, when the blood is examined with a low power, gives to the eye a peculiar impression of disturbance, and will immediately lead the practised observer to look for the presence of spirilla. If he then increases the power, and, still better, if he employs an oil immersion-lens with Abbe's condenser and a small

* In opposition to this view, see *Naunyn*, Centralbl. für Bacter. und Parasitenk., v. 376, 1888.

diaphragm, the spirilla come clearly into view. These bodies are extremely sensitive to reagents of all kinds. Even the addition of distilled water will cause them to disappear.

The number of spirilla which are to be seen together in a specimen of blood varies greatly, and often bears no relation to the intensity of the fever.

If the blood be examined in the intervals of the disease, provided another paroxysm is impending, it displays peculiar refractive bodies resembling diplococci, which are especially numerous when the paroxysm sets in; and just as it begins, they even seemed to the author in certain cases to grow out, as it were, into short thick rods, from which the spirilla were finally evolved. Before those of the author, similar observations had been made by *Sarnow*; ²⁰⁰ and pending further confirmation, it seems probable that these bodies are the spores of spirilla, which have so long been sought for. In addition to spirilla, the blood, especially after a paroxysm, contains pigment particles (melanin), either free or incorporated with the leucocytes.

Since both the spirilla and the forms (?spores) just mentioned have been met with as yet only in the blood of persons suffering from relapsing fever,* their great importance as a clinical test is sufficiently apparent.

It should be mentioned, however, that forms very similar to the spirillum of relapsing fever are met with in the blood of malarial patients (see p. 65) (*v. Jaksch* and *Canalis*). In such cases the two diseases may be distinguished by attending to the other characters of the blood. Moreover, when relapsing fever supervenes on an attack of ague, the spirilla assume a modified shape, as it would appear from a very interesting observation by *Karlinski*.²⁰¹

According to *Sacharoff*,²⁰² the masses of protoplasm which have been seen in the blood of relapsing fever constitute the specific haematozoon of that disease. He believes that these develop in the red corpuscles, and that portions of their nuclei give origin to the spirochæte bodies. The writer in question attempts to account in this way for certain resemblances between ague and relapsing fever. The effort to cultivate these micro-organisms outside the blood has not yet succeeded; but the disease can be transmitted to monkeys by means of the blood (*Carter, Koch*). *Pasternatzkij*²⁰³ has observed the parasite to survive for a day within the alimentary tract of the leech. In Russia the opinion prevails amongst medical men that relapsing fever is commonly propagated by insects (flies).²⁰⁴

For the purposes of research an unaltered specimen of blood serves best, but dried preparations can also be made and stained most appropriately with fuchsin.

* Bodies resembling them are found in the buccal secretion (*vide chapter on the buccal fluids*).

Günther²⁰⁵ has recently advocated the following plan:—The cover-glasses, prepared in the usual manner, are immersed for ten seconds in a 5 per cent. solution of acetic acid, to destroy the colour of the red corpuscles. The acetic acid is then removed as far as possible by blowing upon the cover-glass, and its last traces neutralised by holding the preparation surface over an open flask of strong ammonia solution (shaken up previously). The preparation is next stained in the Ehrlich-Weyert aniline-water gentian-violet solution, and the staining fluid washed off. It is then mounted in Canada balsam or xylol before it is inspected. This method is a very good one for the examination of the blood for micro-organisms in general.

3. **Tubercle Bacillus.**—This was first described by Weichselbaum²⁰⁶ as occurring post-mortem in a case of miliary tuberculosis. To one of his pupils named Meisels²⁰⁷ belongs the credit of its first discovery in the same connection during life. Since then much knowledge on the



FIG. 25. Tubercle Bacilli in the Blood, from a preparation by Professor Weichselbaum (eyepiece III, objective Zeiss $\frac{1}{2}$, homogeneous immersion; Abbe's mirror, open condenser).

subject has been derived from the labours of various observers (*Lustig, Sticker, Doutrelepont, Rüttmeyer*).²⁰⁸

The assertion of Liebmann²⁰⁹ that tubercle bacilli occur in the blood of patients who have undergone tuberculin injection is not confirmed by Ehrlich, Guttman,²¹⁰ Hamerle,²¹¹ and others (Kossel).

The number of tubercle bacilli found together is commonly small. Indeed, when present, they are often so few as to baffle the most careful investigation. It rarely happens that a specimen of blood is found so rich in bacilli as that represented in fig. 25; but, when detected, they leave no doubt of the existence of general miliary tuberculosis.

The mode of research is the same as that detailed at p. 126 for the examination of the tubercle-bacilli of the sputum, and the cover glass is prepared in the manner already described.

4. **Bacillus of Glanders.**—The bacillus of glanders was first discovered by Loeffler²¹² and Schütz,²¹³ and its association with the disease has been

confirmed by *Israel*²¹⁴ and *Weichselbaum*.²¹⁵ The bacilli are rod-like bodies, 2–3 μ long and 0.3–0.4 μ broad, often carrying a spore at the extremity. They are found in the farcy buds and ulceration of glanders, and also in the blood of persons suffering from that disease. Fig. 26 represents the bacilli seen in a specimen of blood taken from a case of glanders in the Vienna General Hospital. They may be shown in the blood by means of the dried preparation, and they are best stained by Löffler's method for detecting these fungi. (See Chapter VIII.)

5. Bacillus of Typhoid Fever.—Bacilli have of late years repeatedly been found in the blood of typhoid patients, and they are doubtless the exciting cause of the disease. *Meisels, Rütimeyer, and Neuhauss*²¹⁶ experimented upon the blood taken from the roseolar spots of typhoid patients, and succeeded in cultivating bacilli from it. Recent researches, as those of *Janowsky*,²¹⁷ show that the bacillus is very rarely present in the blood during life, and that its detection should not be numbered amongst the resources of diagnosis. In support of this, and as a result

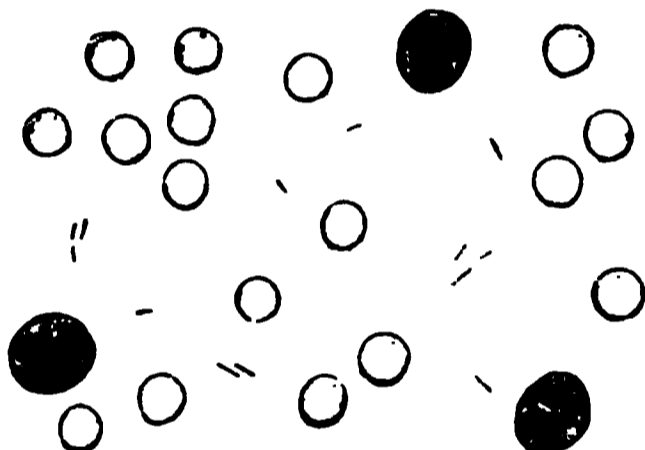


FIG. 26.—Bacilli of Glanders in Human Blood, from a preparation by Dr. Kolisko (eye-piece V., objective Zeiss $\frac{1}{2}$, homogeneous immersion; Abbe's mirror; open condenser).

of very numerous observations of the blood of typhoid patients in every stage of the malady, the author can state that neither microscopically nor by process of cultivation has he been able to determine the presence of bacilli. Further information on this subject will be found in the chapter on *Faeces*.

6. Staphylococci and Streptococci.—*v. Noorden*²¹⁸ recently found streptococci in blood taken from the body of a woman who had died of erysipelas. They exhibited in their cultivation-properties the closest resemblance to the well-known streptococci of *Fehleisen* and *Rosenbach*.

In like manner, *Orthenberger*,²¹⁹ by applying Weigert's method to the examination of the blood after death, has shown the presence in it of pneumococci in six cases of uncomplicated lobar pneumonia.

In a series of cases in which the author examined the blood of pneumonia patients, employing sterilised human blood-serum as the nutrient substance, he failed altogether to obtain a cultivation of cocci.

Some years ago *Klebs* ascertained that micro-organisms existed in the blood and endocardial vegetations of persons with endocarditis. In a case of congenital heart-disease with endocarditis, *Sünger*²²⁰ has cultivated micro-organisms obtained from the blood during life. An instructive communication lately published by *Weichselbaum*²²¹ has also shown that cultivations of cocci can be obtained from the blood during life in cases of endocarditis. That other agencies are productive of endocarditis is sufficiently proved by the observations of *Leyden*, *Howard*, and *Chiari*,²²² amongst these agencies being the bacillus of diphtheria, gonococci, and other micro-organisms. The author has found micro-organisms in the blood of acute rheumatism, and made cultivations of them outside the body. These were most commonly cocci. It is notable that the infection of animals with these micro-organisms was often unattended by any result, the inference being that they had already expended their virulence (*Sahli*).²²³ [An alternative view, founded upon clinical experience, and especially upon the observation of hereditary influence, is that the micro-organisms associated with endocarditis are not by themselves productive of that condition, but require for their existence or operation a peculiar condition of the blood (*Cagney*),²²⁴ and this may be connected with the arthritic disorders, some of which are called rheumatoid arthritis.] The blood of patients in a severe attack of puerperal fever has repeatedly been found to exhibit micro-organisms of this class (*A. v. Rosthorn*).²²⁵ The researches of *A. v. Eiselsberg*, *Levy*, *Brunner*, and *A. Huber* have shown remarkable results.²²⁶ In the author's clinic *Ott*²²⁷ has found cocci in the blood of puerperal septicaemia, and in another case of obscure sepsis the diagnosis was made by the discovery of cocci, and confirmed post-mortem, when it was found to proceed from an abscess of the verminiform appendix.

Sottman,²²⁸ by using an appropriate method, was able to demonstrate cocci in the blood of every case of septicaemia examined. *Jakowski*²²⁹ found various staphylococci in the blood in each of seven cases of phthisis with high temperatures. *Cerny* and *Moser*²³⁰ observed micro-organisms in the blood in twelve cases of children with gastroenteritis. *Brunner*²³¹ detected staphylococci in the blood of a patient suffering from osteomyelitis. It should be mentioned here that *Verdelli*²³² professes to have repeatedly seen micro-organisms in the blood of leukæmia and pseudo-leukæmia, but the most careful researches made by the author in eight cases of leukæmia were without result.

[*Streptococcus Pyogenes*.—A chain-forming micrococcus was described by *Ogston* as occurring in abscesses. It was afterwards submitted to Koch's cultivation processes by *Rosenbach*, who gave it the name of *Streptococcus pyogenes*, and it is now known to be associated with septic processes of all kinds, whether in men or animals. It is apparently the same as *Fehleisen*'s *S. erysipelatis* and *Löffler*'s

Diphtheriacoccus. It has been obtained from the pus of pyæmic abscesses and empyema, from the tissue-fluids in spreading gangrene, from the vesicles and pustules of small-pox, and in contaminated calf's lymph. It occurs in the blood-vessels in certain cases of diphtheria, scarlet fever, puerperal septicæmia, measles, and typhoid; but it must be regarded only as a secondary result of those affections. It may gain admission to the system in any disease attended with lesions to the skin or mucous membrane, setting up destructive processes, when the resistance of the tissues is greatly impaired by the working of a special virus (*Crookshank*).²³³

Streptococcus of Scarlatina.—*Dr. Klein* has succeeded in separating a micro-organism obtained from the blood of scarlatina patients, which he regards as the contagium, virus, or actual cause of that disease. When cultivated by inoculating the surface of nutrient gelatine with the post-mortem blood, the micro-organisms bear a close resemblance to the *Streptococcus pyogenes*. Klein is of opinion that the two are sufficiently distinguished, both morphologically and by their pathogenic effects upon animals. The experiments upon which he relies, however, are questioned by *Crookshank*,²³³ who maintains the identity of the micro-organisms concerned. A *Streptococcus* which Klein believes to be the same as that of scarlatina was found in the blood of cows in an outbreak of disease amongst those animals at Hendon; and this, when inoculated upon calves, produced in them the post-mortem appearances of scarlatina. Moreover, the cow-disease alluded to is thought to have caused an epidemic of scarlet fever amongst men through the medium of the milk.^{234]}

7. **Micro-Organism of the Blood in Hydrophobia.**—*Bareggii*²³⁵ found constantly present in the blood in hydrophobia a micro-organism which stained with methylene-blue. Placed on slices of potato at 25°-27° C., it developed in the course of forty-eight hours in flattened hemispherical cultivations, of a colour ranging between whitish-grey, yellow, and citron. In test-tube cultivations (see the chapter on *Bacteriological Methods*) it behaved like the bacillus of Asiatic cholera. Further research is needed to establish a connection between this micro-organism and hydrophobia.

8. **Bacillus of Tetanus.**—Many authorities²³⁶ are of opinion that tetanus is an infectious disease, produced by the agency of the "bristle-bacillus" of *Nikolaier*. According to the observations of the latter investigator, the bacilli in question are somewhat longer and thicker than those of mouse-septicæmia. They sometimes form in threads, commonly in irregular heaps, and occasionally produce spores. The bacilli themselves, or their spores, have been found in the blood (?) (*Hochsinger*). They stain readily in dry cover-glass preparations, and may be cultivated outside the system. The fact may be mentioned here that from cultivations of the bacillus *Brieger*²³⁷ has isolated several ptonaines—tetanin, tetanotoxin, and spasmotoxin—with which he has induced tetanic poisoning symptoms in animals; and more recently the same observer has obtained similar poisons from the blood of persons who had died of tetanus.

*Nissen*²³⁸ also has experimentally demonstrated the presence of toxin in the blood in cases of tetanus. [*A. Bruschettini*²³⁹ has lately succeeded in causing fatal tetanus in rats and rabbits by injecting the urine of

patients suffering from that disease.] The researches of *Kitasato*²⁴⁰ in Koch's laboratory have definitely established the fact that the bacillus is the specific cause of the disease. It is anaerobic, develops in pus and forms spores there; but it is occasionally observed in the form of rods free from spores, if the pus be examined in an early stage.²⁴¹

As *Nissen* has shown that the blood of tetanus contains a substance capable of producing the disease in mice, this fact may be utilised in doubtful cases where bacteriological processes are insufficient to clear up the diagnosis. *Walko*²⁴² has recorded a case of puerperal tetanus in which he adopted this course.

9. Influenza-Bacillus.—Against the statement of *Canon*,²⁴³ who thought he had found a bacillus in the blood of influenza patients, is the experience of other observers (*Pfeiffer, v. Jaksch*).²⁴⁴ The matter requires further investigation. (See Chapter IV.)

10. Bacterium Coli Commune.—The experience of recent years has tended greatly to extend the rôle of this parasite, and there is no longer doubt that it occurs in the blood as well as elsewhere.²⁴⁵ In connection with the faeces, where it is innocuous; with the urine, wherein its presence is always marked by profound constitutional disturbance; with peritonitis and infection from wounds,²⁴⁶ it will be separately noticed in the appropriate places. *Canon, Pielicke, and Czajkowski*²⁴⁷ attest the presence of various bacilli in the blood in measles and other infectious diseases.

11. Plague Bacillus.—The widespread occurrence of the plague since its descent from the Great Plain of China to the sea, coupled with the circumstance that Europe also is decidedly in danger of becoming infested with this disease, seems to justify mention of the discovery of *Kitasato*²⁴⁸ and *Yersin*.²⁴⁹ These workers regard as the instigator of the plague certain micro-organisms (bacilli) which they almost invariably detected in the blood of the persons and animals infected with this disease. Observations made by *Kitasato* show that plague bacilli are found in the blood of convalescents after the disease has passed away, and that hence the plague can also be accurately diagnosed after it has run its course. It should likewise be mentioned that these micro-organisms have also been found in the excreta, in the bubonic pus and in the urine of sufferers from this plague; their detection in the blood, however, constitutes the most important point in the diagnosis of the disease.

According to the observations of other writers, of whom only *R. Abel*²⁵⁰ and *Klein*²⁵¹ will be cited here, the plague bacillus exhibits an unusual degree of polymorphism. Numerous plump rods, with rounded ends and incapable of standing the Gram method of staining, are met with in the blood and glands, the bacilli being of variable dimensions—often as broad as long, whereas in other instances they attain to considerable

length, and may also take the form of chains resembling streptococci. Spindle-shaped cells are also met with. According to *Klein*,²⁵¹ the plague bacillus thrives on nutrient gelatin composed of beef broth with 10 per cent. of gelatin and 1 per cent. each of common salt and peptone. It also grows on all the usual culture media. At the end of twenty-four hours the cultures appear as small grey spots with rounded edges; and according to *Klein* the young colonies exhibit a marked resemblance to colonies of *Proteus vulgaris*. The same authority regards as characteristic of the plague bacillus the appearance of occasional atypical thread colonies. The animal experiment is important. Injection into the peritoneum of the porpoise gives rise to an exudation in which the characteristic chains of the plague bacillus can be detected.

The bacillus is ciliated but non-motile (*Meroyn Gordon*²⁵²); nothing is as yet known concerning its power of forming spores. It can be stained with the usual aniline colours.

From existing data, this dangerous foe to the human race morphologically resembles the species of *Proteus*. The detection of the plump bacilli in the blood is of prime importance in the diagnosis of the disease.

B. Animal Parasites (Hæmatozoa).

1. **Protozoa.**—Under this heading are to be described the micro-organisms of malaria—parasites of very great interest, which occur in the two principal varieties of *Hæmamœba malariae* and *Laverania malariae*.

Historical.—*Klebs*²⁵³ and *Tommasi-Crudeli* have described a bacillus which is found in the soil of the Campagna, and which they regard as the specific cause of malaria.

To *Lareran*,²⁵⁴ who, in 1880, observed flagellated organisms in the blood of malaria patients, belongs the credit of the first step towards the ultimate pathology of this disease; but his discovery was too indefinite to lend itself to any clinical purpose, and *Marchiafava* and *Celli*²⁵⁵ claim the honour of having first distinguished and described the hæmatozoon, which is the most abundant, and, clinically, the most important, of this group of blood-parasites. These observers detected amœboid bodies within the red corpuscles in the blood of persons infected with malaria. The amœboid bodies (plasmodia) commonly contained granules and particles of black pigment. They stain with methylene-blue. They have not yet been cultivated outside the body,²⁵⁶ but *Marchiafava* and *Celli*, and also *Gerhardt*,²⁵⁷ by inoculation and intravenous injection with malarial blood containing them, has succeeded in communicating the disease to other individuals, and the infected blood in its turn exhibited the plasmodia. The statements of these authors are borne out by others.²⁵⁸ The author has lately made an attempt of this kind without result. He injected blood containing the plasmodium, and derived from a case of tertian ague, subcutaneously into a patient with carcinoma of the stomach.

[*Fenton Evans* has succeeding in obtaining pure cultivations of the hæmatozoon outside the body by treating the nutrient media with living blood—i.e. blood taken before *rigor mortis* has set in. These cultivations, when inoculated upon animals, produced a fatal disease, of which the symptoms were intermittent,

though not the same as those of classical intermittent fever. He believed also that by altering the chemical composition of the media the haematozoon attained a higher organisation. This statement is questioned by *Dyer*.^{259]}

Metschnikoff proposes to name the parasite *Hæmatophyllum malariae*. *W. Oader* has examined the blood in seventy cases of malarial disease, and found it present in all. It would appear from his researches, however, that the organisms concerned assume a greater variety of forms than was previously supposed.

Councilman,²⁶⁰ again, has recently supported the views of *Laveran*, *Marchiafava*, and *Celli*. He describes several forms of the parasite, and opposes the doctrine of *Mosso*,²⁶¹ who sought to prove by his experiments that the plasmodium resolves itself into degeneration types of the red blood-corpuscles ; and maintains that similar appearances can be found in the blood apart from malaria. The author's experience enables him to agree with *Mosso*, and also with *Maragliano* and *Castellino*,²⁶² in attesting the occurrence of similar changes in the shape of the blood-corpuscles independently of the presence of the plasmodia ; but he holds that the latter are sufficiently characteristic in appearance. *Tommasi-Crudeli*,²⁶³ on the other hand, regards the forms in question rather as the result than as the cause of malarial disease, believing them to be the product of the degeneration of the red blood-corpuscles, and further maintains that the bacillus discovered by *Klebs* and himself is the true cause of the disease. This opinion is shared by *Schiaruzzi*,²⁶⁴ who still more recently has succeeded in cultivating the micro-organism outside the system. The theory of *Danilevsky*²⁶⁵ remains to be mentioned. That author supposes that the bodies under discussion are identical with the haematozoa observed in the blood of many birds. In a recent communication he suggests for them the name of *Polymitus malariae*, and from his researches, as well as those of *Grassi*, *Feletti*, *Celli*, and *Sanfelice*,²⁶⁶ it appears probable that birds are attacked by the same parasite which is associated with malaria in man.

Finally, *L. Pfeiffer*²⁶⁷ has seen similar forms in the blood in scarlatina and after vaccination.

Personal observations, no less than the study of what has been written on the subject, have satisfied the author that *the blood in malaria contains specific organisms*, as *Laveran* first pointed out, and that these specific organisms are the determining cause of the disease. The subject is thus invested with great importance from the point of view of diagnosis ; but it must be admitted that the micro-organisms in question display a great variety of form, and many details of the life-history of this interesting parasite remain yet to be discovered. For our purpose, it will suffice to mention those particulars which at present admit of clinical application. The following description is founded on the statements of *Laveran*, *Marchiafava* and *Celli*, *Golgi*, *Celli* and *Guarnieri*, *Grassi* and *Feletti*, *Canalis*, *R. Paltauf*, *Quincke*, *Dolega*, *Plehn*, *Chenzinsky*, *Rosenbach* and *Rosin*, controlled by the author's own observations.²⁶⁸ Some of the figures are borrowed from these writers.

The earlier researches of *Laveran*, which are so far confirmed by the Italian observers, have established two facts—first, that the parasite of malaria assumes a variety of forms in different cases, even within the same locality ; and secondly, that its form tends especially to vary with

the locality and with the clinical type of the disease. Hence the discrepancies in the statements made by competent authorities, as by *Laveran* on the one hand and *Golgi* on the other, the former having had occasion to see only or chiefly the spherical and crescentic bodies presently to be noticed, while the amoeboid parasite seldom came under his observation.²⁶⁹

We have it on the authority of *Marchiafava*, *Celli*, and *Canalis*, and especially of *Golgi* and *Canalis*, that the haematozoon of malaria occurs in three principal forms, each having a special relation to the



FIG. 27.—Parasite of Tertian Ague, as seen in the blood some hours after a paroxysm
(partly copied from other observers).²⁷⁵

different clinical types of the disease—tertian and quartan ague, the anomalous varieties, remitting and intermitting attacks, and cases of fever with short periods of apyrexia (*Febris perniciosa algida*) (*Marchiafava* and *Celli*), and that the development of the organism within the system is intimately associated with the production of the various sets of symptoms.

[*Thayer* and *Hewetson*,²⁷⁰ in an exhaustive résumé of the literature of the subject, express a similar opinion. *Marshall* and *Thin*,²⁷¹ who investigated the appearances presented by the blood of malaria patients met with at Huelva, in Spain, and who carefully collated the clinical symptoms



FIG. 28.—Progressive Endoglobular Development of the Parasite during the Apyrexia Interval (in part copied from *Golgi*).²⁷⁵

displayed by these patients, also maintain that the varieties of the parasite are associated with the varieties of the disease. *Curnow*²⁷² opposes this view. *Hehir*²⁷³ has very carefully investigated the blood of malaria patients in India, and the results, which he has lately published, agree very closely with the statements of *Laveran* and the Italian observers. He found that there were constantly present in such blood eight different forms, including the phagocytes, which may be observed to take an active part in the morbid process. The micro-organisms are, in his opinion, all varieties of a single polymorphous haematozoon, and in general they

correspond to Laveran's description. The chief interest attaches to a micro-organism with flagellated rapidly-vibrating processes, which was very partial in its distribution through the body, was always most abundant in the pyrexial period, and the vitality of which was controlled by quinine. There was also found a spherical body with 3-6 well-formed cilia-like processes. This is the *Hæmatomonas maliariæ stellata*. *Lawrie*²⁷⁴ examined the blood of malaria patients in India and failed to find the plasmodium. He regards the bodies observed by Laveran as the products of blood destruction in the spleen.]

1. The Parasite of Tertian Ague.—A few hours after the cessation of the febrile paroxysm the blood may be seen to contain very small movable bodies of a pale colour, and carrying from one to three extremely delicate and pigmented thread-like processes (ekto-globular parasite, fig. 27 on the left). *Plehn* and the author have observed similar bodies during the fever-free period.

The parasite, it is supposed, then invades, or has already invaded, the red corpuscles (fig. 27, left), where it appears as an actively-moving body,

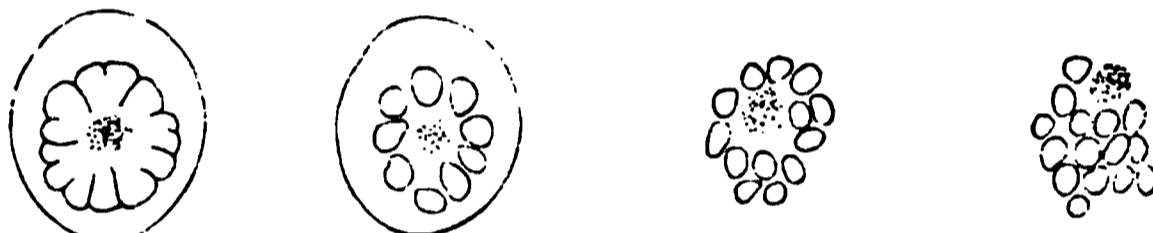


FIG. 29.—Parasite of Tertian Ague, at or just before the commencement of the attack (copied from *Golgi*).²⁷⁵ The figure shows the various segmentation forms of the parasite.

already, it may be, outlined, as it were, with a wall of melanin granules. Within the corpuscle it grows in size, and as it does so, the former becomes progressively despoiled of its haemoglobin. In this way the parasite develops into a large freely-moving, deeply-pigmented mass of protoplasm (*Hæmamœba*, see p. 57). The changes just described occur in the fever-free period within twenty-four hours from the termination of a paroxysm.

While the corpuscles which have been attacked rapidly lose their colour (fig. 28), the newly-formed melanin accumulates towards their centre, and the parasite comes to occupy the entire contents of the corpuscle. The parasite then undergoes segmentation, and from this result the various forms described by *Golgi*, and represented in fig. 28. These segmentation forms will be noticed at greater length presently.

The invaded corpuscle is disintegrated. Two more days are required for this endoglobular development of the parasite. This terminates the apyrexial period. A new generation of the parasite has matured, and its presence in the blood determines a new paroxysm of fever.

The development of the parasite is often completed before the paroxysm begins.

2. The Parasite of Quartan Ague.—The course of quartan ague is marked by a development similar to that just described (*Golgi*).²⁷⁶ The endoglobular growth takes place in this case also during the fever-free period. Its earlier phase is quite the same as that of tertian, only the corpuscles are less rapidly decolorised, and the melanin granules formed are of larger size.

The chief difference, however, is in the method of segmentation, the resulting segments being much fewer in quartan than in tertian ague (fig. 30). In the latter they may be as many as from 15–20 for each plasmodium, whereas in the former they range from 6 to 12. Moreover,

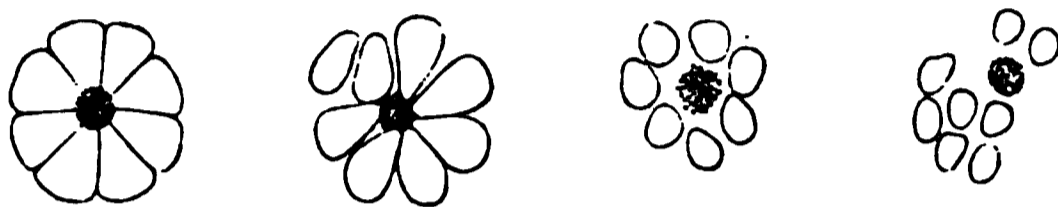


FIG. 30.—Parasite of Quartan Ague. Various segmentation forms (from *Golgi*).²⁷⁶

the process of segmentation itself takes place in a more uniform manner in quartan fever. The parasite requires three days for its development. *Golgi* asserts that in quotidian fever the blood contains at once three broods of the parasite of quartan, which mature successively at intervals of one day.

3. The Parasites of Acyclical and Anomalous Forms of Ague.—For a knowledge of these organisms we are indebted to *Celli* and *Marchiafava*,²⁷⁷ and especially to *Canalis*. *Marchiafava's* observations were made with the blood of patients suffering from the acyclic forms of intermittent fever which prevail at Rome in the summer,



FIG. 31.—Parasites of Acyclical Intermittent Fever (from *Celli* and *Marchiafava*).

autumn, and winter months. In Roman fever, just before the paroxysm and at the end of the apyrexial period, the blood contains small annular plasmodia, holding in their central part a mass of haemoglobin or pigment granules; also minute moving amoeboid bodies with a sinuous outline, and large round stationary forms which are almost colourless, and which present at the centre or periphery a circular spot of pigment. According to *F. Plehn*,²⁷⁸ a small parasite (which has no pigment) is found in the Cameroons, on the West Coast of Africa, in the blood of individuals attacked by the so-called "black-water fever." Unlike the parasites of tertian and quartan ague, the plasmodia of the æstivo-autumnal

Roman fever are devoid of pigment, and retain their motility for a long time (*Celli* and *Marchiafava*).

[*Kanthack*²⁷⁹ has described a micro-organism derived from a case of malaria acquired in Ashanti. It resembled the *Hæmamœba præcox*. It was ovoid, with a curved axis, and centrally pigmented.]

In the varieties of Roman fever just noticed are often to be found the crescentic and sickle-shaped bodies which were first described by *Laveran*. According to *Celli* and *Guarnieri* the following forms may be distinguished: Crescented or sickle-shaped bodies, others of a boat, or spindle-shape, and again others with an oval or circular outline, and provided with flagella (fig. 32).

Grassi and *Feletti* have urged that the name of *Hæmamœba malariae* should be appropriated to those forms of the plasmodium which have been described as belonging to the recognised typical varieties of ague, while the term *Laverania malariae* is reserved for the crescentic micro-

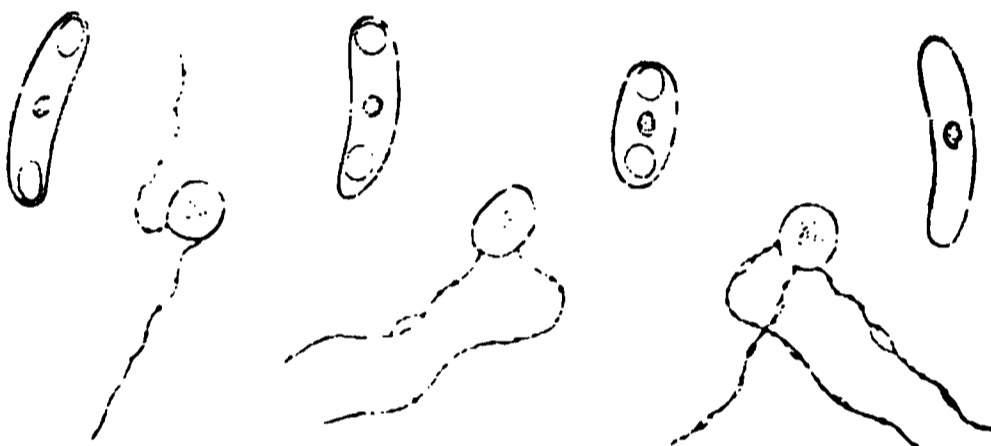


FIG 32.—Crescentic and Falciform Bodies, and Free Organisms carrying Flagella (*Celli* and *Guarnieri*²⁸⁰).

organism. These writers are of opinion that the latter is associated with recurring attacks and the malarial cachexia. *Golgi* holds the view that the *Laverania* occurs in cases of malarial fever, where the attacks are separated by long and indeterminate intervals of immunity.

[According to *Curnow*,²⁸¹ the circular or amoeboid plasmodium occurs in the blood in all forms of malaria, whatever the type, as they present themselves at the Seamen's Hospital. He rejects the conclusion that special developments are specifically connected with quotidian, tertian, and quartan ague. The micro-organism of quotidian fever has been described by *Marchiafava* as of two kinds, the pigmented and the hyaline; in this he is approved by *Grassi* and *Feletti*, while *Golgi* denies the existence of the unpigmented parasite. The nature of these forms and their relation to the quotidian type of fever are far from determined, one reason for this being that the clinical character of the disease in which they have been found is not clearly established.

According to *Manson*,²⁸² the rosette-shaped body is the final and mature form of the parasite. He regards the petals as spores which fall off and assume amoeboid movements. They penetrate within a blood-corpuscle and grow at the expense of the haemoglobin, which they convert into pigment. Later this pigment accumulates at the centre of the plasmodium, which now assumes the circular form ; and from this, again, the rosette results by a process of segmentation. The petals, as spores, survive. The central body, with its pigment, is removed by phagocytes, and deposited in the spleen and other organs. Much interest attaches to the crescentic form of the parasite. Of this, again, *Curnow* asserts that, in his experience, it is to be met with in every form of malaria. Most writers are agreed that it is associated chiefly with cases of chronic and persistent disease. It is of all forms the most resistant to the influence of quinine (*Bacelli*, *Curnow*). Its origin is a matter of dispute. *Manson*²⁸² believes that it is the centrally pigmented plasmodium distending a corpuscle and itself distorted in consequence. *Grassi* and *Feletti* describe its sporulation and reproduction ; but there is also evidence that it may result from the circular variety. *Thayer* and *Heinetson* regard it as a development of the smaller hyaline parasite. The crescent especially, but also the tertian and quartan parasite, become flagellated some time after removal from the body. They can then be seen moving actively in the microscopic slide. The development of flagelli has been regarded by many observers as a moribund or degenerative phenomenon (*Marchiafava*, *Grassi*, *Begnami*, *Blanchard*). *Grassi* and *Feletti* assert that the nucleus takes no part in the formation of the flagella, and suppose that the movements are those of dying protoplasm, such as are manifested by the amoebæ. *Sakharoff*,²⁸³ on the other hand, asserts the participation of the nucleus in the process of flagella formation. *Golgi* thinks that the flagellated form may be a transitional state of the parasite ; and *Manson*,²⁸⁴ arguing from analogy with what he has observed in the course of development of *Filaria*, maintains that the flagella are an extra-corporeal organ by means of which the parasite is fitted to maintain itself in a new host. *Manson's* hypothetical host is (as in the case of *Filaria*) the mosquito, and by that insect this observer supposes that man is again infected through the medium of water and air. *Lareran* and *Pfeiffer* had similarly attributed to mosquitoes an important rôle in the dissemination of the plasmodium, while *Grassi* and *Feletti* have combated this notion. The observations of *Marshall* and *Thin* go to show that, in the province of Huelva at all events, telluric conditions are the prominent factor in infection ; that the plasmodium is not usually conveyed by water ; and that there is no evidence of the mosquito being an intermediate host.

*Manson*²⁸² rejects *Golgi's* view that the periodicity of attacks depends

upon the maturation of successive crops of the plasmodium, and refers this rather to a physiological rhythm in the resistive power of the body.

Larrie,²⁸⁵ who denies the existence of a plasmodium in the blood of malaria, attributes the disease exclusively to climatic conditions. It is essentially an inflammatory affection of the spleen, and the bodies which have been mistaken for a parasite are abortive or defective corpuscles due to this.}

Canalis has directed his attention particularly to the study of the parasites which are to be found in cases where the successive febrile attacks are separated by more or less considerable intervals, and which, for the most part, result in the malarial cachexia. In this connection

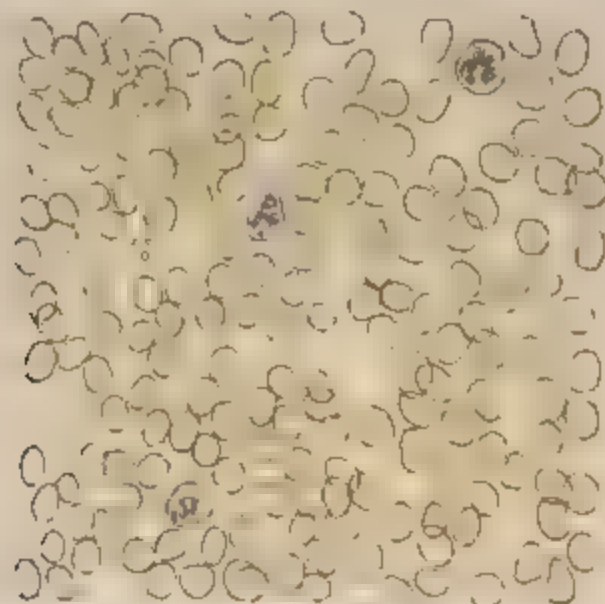


FIG. 32.—Tertian Ague; the blood at the commencement of the attack. The figure is composed so as to represent the appearances observed in several cases. Zeiss's compensation eye piece 4, apochromatic objective 2 mm.; apert. 1.40, homogeneous immersion; Abbe's mirror, and medium diaphragm.

he has described the life history of two varieties of *Laverania*, and he also finds that the period of maturation of one brood of the parasite coincides with the development of fever.

Without entering into details, the author would observe that *Canalis'* description applies to certain forms of the haematozoon which he himself has observed in a case of reduplicated quartan ague with a very irregular course, and in certain other particulars the observation is worthy of notice. Besides the usual amoeboid forms, there were to be seen a remarkable number of pale, homogeneous red blood-corpuscles, and immediately after the febrile period—a double paroxysm lasting for twelve hours—the blood contained free particles of protoplasm, enclosing fine granular pigment and having attached long and well-formed flagella. There were also present small round bodies segmented in the middle and provided with long, thick, freely-moving flagella, upon which black particles were scattered. (See fig. 32.)

The most curious fact noticed was the presence, mostly towards the outer boundary of the preparation, of spirally twisted bodies resembling the spirillum of relapsing fever, but thicker and longer than the latter, and further distinguished by their outline being broken by very minute pigment particles. These bodies were actively mobile in the direction of their long axis. They never made their appearance until some hours after the preparation had been mounted.²⁸⁸

Clinically, as has been said, the case was one of reduplicated quartan ague, and pathologically it was found that distinct generations of the parasite matured at separate intervals, their maturity corresponding with the access of fever.

Besides the parasites described here, the blood in malaria contains pigmented leucocytes; these, however, are not characteristic of the disease, occurring as they do in other febrile states, as, for instance, in relapsing fever.

As the result of the foregoing description, it will be seen that the parasite of malarial fever exhibits a singular variety of form.

Notwithstanding, there are certain well-established facts available for the diagnosis of the disease as it presents itself in European climates. *An examination of the blood at the outset, or during the period of fever, will suffice to establish the nature of the disease in cases of tertian ague. A number of the red corpuscles will be seen to be of a remarkably pale colour, and within some of these pale corpuscles will be found freely-moving colourless bodies, containing a fine granular pigment. Some of the red corpuscles are almost quite decolorised, and within these the process of segmentation of the Hæmamæba into 15–20 parts may be more or less clearly distinguished. On the other hand, where this process can be observed, and where the segments are fewer—6 to 8 in number—and result in the characteristic marigold arrangement, the diagnosis of quartan fever may be made with equal certainty.*

In view of the great importance of this subject, it is deemed advisable to illustrate the foregoing remarks by a figure representing a specimen of blood taken from a case of tertian fever at the commencement of the paroxysm. The pale corpuscles contain the plasmodia (fig. 33).

This figure is drawn from appearances actually observed by the author in three cases of tertian fever, and it will perhaps have an additional value as affording a means of comparison with figs. 27–29, which were partly borrowed from other sources, and in part merely diagrammatic.

In cases where the Hæmamæba is found in the blood, together with the forms of Laverania just described, the condition may be taken to be one of atypical or anomalous intermittent fever.

It is unnecessary to dwell upon the great importance of these researches. By their aid it has become for the first time possible to distinguish malaria with absolute certainty by the result of an examination of the blood, and we have thus acquired an invaluable means of discriminating other affections which closely resemble it, as obscure sepsis,

and certain cases of endocarditis and tuberculosia.²⁸⁷ Mention should not be omitted that, of late years, *Golgi's* views have been combated from different sides. Nevertheless, although the author admits the justice of several of these objections, his own experience in regard to the behaviour of the blood in malaria convincingly shows that none of them destroys the diagnostic importance of these forms.

4. Methods of Examining the Blood for the Parasites of Malaria.—No more need be said to impress upon the reader the expediency of being able to recognise the principal varieties of this interesting parasite. For the purpose is needed an oil-immersion lens and a

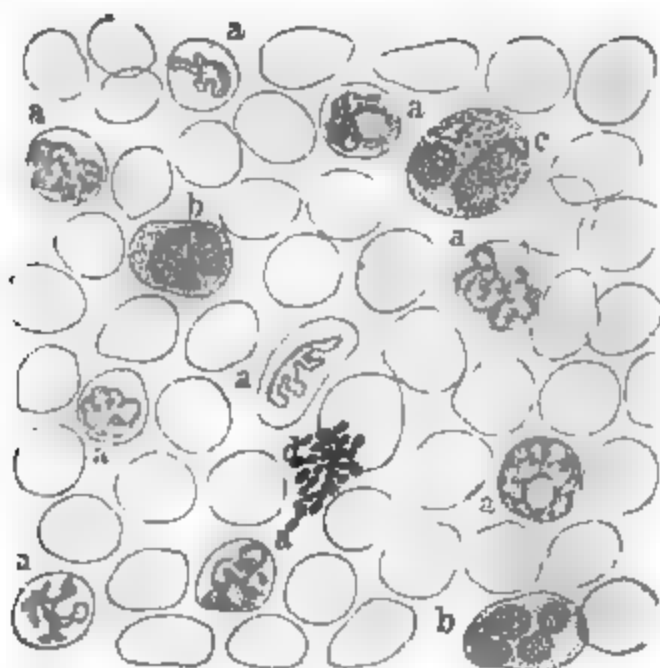


FIG. 34.—The Blood in Tertian Ague at the Commencement of an Attack. Stained by Alderhof's method; a. Plasmodia; b. Leucocytes; c. Eosinophil-leucocytes; d. Blood-platelets (compensation eye-piece VIII., apochromatic objective 2 mm., apert. : 40, homogeneous immersion, Abbe's mirror and open condenser.)

moderately wide aperture of the diaphragm, or, still better, an apochromatic objective—as Zeiss's apochromatic objective $\frac{2.0}{1.40}$ with compensating eye-piece IV., or Reichert's semi-apochromatic $\frac{1}{12} 18^b$. With such an instrument very little practice will secure proficiency of observation; indeed, the endoglobular pigmented parasite can be discovered quite as readily as, for instance, the spirillum of relapsing fever.

For the purposes of more precise investigation, and especially to avoid the chance of being misled by the process of vacuolation in the red corpuscles which has been noticed at p. 41, it is necessary to resort to staining methods.

To distinguish the parasite of malaria from simple vacuolation it will suffice to smear the under-surface of the slide upon which a preparation is mounted

with some blue fluid, as, for instance, a solution of aniline dye. The vacuoles will then display the same colour as exists elsewhere in the field when not occupied by corpuscles.²⁸⁸

The parasite may be suitably stained in the following way :—

Methylene-blue is dissolved in normal (0.6%) salt solution until the fluid is somewhat deeply coloured. The latter is then filtered, sterilised, and set apart in small quantities in thoroughly sterilised test-tubes. The point of the finger is then carefully cleansed, a drop of the staining fluid applied to it, and through this the finger is pricked with a needle. The flowing blood mixed with the staining fluid is brought in contact with a cover-glass, and this is placed preparation-side downwards upon a slide and examined. The preparation must be spread in a very thin layer, and it is therefore necessary to guard against evaporation. This may be done by sealing the edge of the cover-glass with paraffin-wax. It may be examined first with a fairly wide aperture of the diaphragm, and afterwards with an open condenser and a good oil-immersion lens.

The plasmodia, whether enclosed within the red corpuscles or lying free in the blood, are stained a distinct blue of a light shade, and upon this the pigment particles which they exhibit and the process of development which they undergo may be easily discerned. It must be mentioned that besides the plasmodia some red corpuscles which are free from them may also take the stain ; but with a little attention confusion will not arise from this, since the corpuscles in question are stained uniformly throughout. Instead of normal salt solutions, diluted and sterilised, ascitic fluid may be employed for mixing the methylene-blue solution (*Guarnieri and Celli*).

To mount a permanent preparation, the blood should be dried in a very thin layer, the cover-glass heated for some time in the usual way, and the preparation stained in eosin-methylene-blue solution (*Chenzinsky, Plehn*).²⁸⁹

Plehn's solution is as follows :—

A concentrated watery solution of methylene-blue	60 parts
½% solution of eosin in 75% alcohol	20 parts
Distilled water	40 parts
To this is added 12 drops of a 20% solution of caustic potash.	

The red corpuscles then appear a light red, leucocytes light blue, and their nuclei a deep blue, the eosinophil granules of the leucocytes a deep red ; the parasites of malaria are stained blue. The method yields good results.

The method of *Aldehoff* and *Gabritschevsky* for staining eosinophil cells may also be applied for the detection of these parasites (fig. 34) in the following manner :—

On cover-glasses prepared as laid down at p. 46, the blood is spread

out in a thin layer, and they are then immersed in a concentrated alcoholic solution of eosin * for half-an-hour in the cold, or for 2–3 minutes with heat, removed and washed with distilled water, then again stained by dipping them once or twice into a concentrated watery solution of methylene-blue, and finally well rinsed with distilled water. It is necessary to obtain the blood rapidly, and to conduct the process without undue delay. If this precaution be neglected, the blood-plates which make their appearance (fig. 34 d) may be thought to have something to do with the disease. *R. Paltauf* has directed attention to this as a source of fallacy, and it may possibly explain the remarkable observations of *Hochsinger*.²⁹⁰

It remains to be noticed that *Loeff* and *Pfeiffer*²⁹¹ have discovered in the blood in SMALL-POX certain protozoa, to which they attribute a pathological significance. *M. Löwit*²⁹¹ has published similar observations in cases of leukæmia and anæmia infantum pseudo-leukæmica.

2. Vermes.—Under this heading we have to describe *Distoma hæmatobium* and *Filaria sanguinis hominis*. Both are worms—the



FIG. 35.—*Distoma Hæmatobium*. Male and Female, with Eggs, from a preparation by Dr. Schieß Bey.

first, of the class *Platoda* and order *Trematoda*;²⁹² the second, of the class *Annelida*, order *Nematoda*, family *Filariadæce*.

1. *Distoma Hæmatobium* (fig. 35).—*Bilharz*²⁹³ has found this parasite in the main trunk and branches of the portal vein, in the splenic and mesenteric veins, and in the venous plexuses of the bladder and rectum. The worm infests the North and East Coasts of Africa, and, according to *Brock*,²⁹⁴ is met with also in South Africa. Except in the blood, the eggs are more commonly found than the parasite itself, and this in the liver, the lungs, the bladder, the urethra, the large intestine, and the urine (see chapter on *Urine*), giving rise to diarrhoea, haematuria, and ulceration of the mucous surfaces of these organs. *Distoma hæmatobium* has not yet, so far as we know, been found in the peripheral blood-vessels, and on this account it is seldom seen in the microscopic examination of the blood during life. The parasite is of a white colour. The male and female are distinct individuals, differing in this from the

* Eosin bläulich 22, Bayer, Elberfeld.

other trematode parasites. The former is from 12-14 mm. long, the latter 16-20 mm., and nearly cylindrical in shape. The male is thicker than the female. There are oral and abdominal suckers anteriorly, and the genital opening in either sex lies close behind the latter. On the abdominal aspect of the male is a deep trench with overlapping edges, which begins just behind the abdominal pore, and serves for the reception of the female (*canalis gynecophorus*).

The eggs are slender bodies, about 0.12 mm. long and 0.04 mm. broad, and furnished with a little spike, which projects from the extremity or at the side.

Ringer²⁹³ discovered a new form of this worm at Tamsui in Formosa, and Manson found the eggs of the same species in the bloody sputa of a Chinese who had lived for a long period in Formosa.

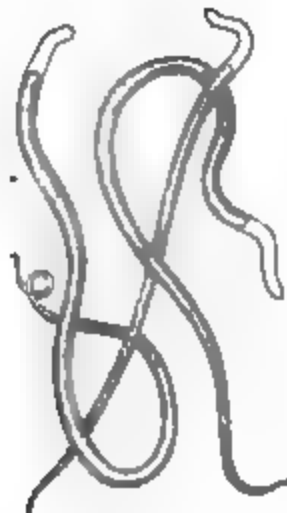


FIG. 36.—*Filaria Sanguinis Hominis* (after Lewis and Leuckart).

2. *Filaria Sanguinis Hominis*.—Dr. Lewis²⁹⁴ of Calcutta was the first to describe it as occurring in the living body.

It is the larva of a filiform worm, *Filaria Bancrofti*, which in the mature state inhabits the human lymphatic system.²⁹⁵ The female is 15 mm. long, and very prolific. The male measures 8 centimetres, and at its hinder end, which is pointed and incurved, carries two spicula of unequal length. The larvae make their way from the lymphatics into the blood stream. The *Filaria sanguinis hominis*, the larva in question, exists in great quantity in the blood. It is 0.007-0.011 mm. in breadth and 0.27-0.34 mm. long. It has a short rounded head, with a tongue-like appendage, and a long pointed tail. From the hinder extremity a ribbon-like process projects, and when viewed with a high power of the microscope, this, like the cephalic appendage, is seen to be the end of a closed sac in which the animal can coil or extend itself. This envelope is entirely structureless, but the contained parasite is seen under a very

high power to be transversely striated and very granular. In the blood the animal exhibits for hours at a time the liveliest movements.²⁹⁸ At first it seems to be homogeneous and transparent, but after some hours its motion ceases, it assumes a darker tint, and the granular contents of its body are easily discernible.

The parasite is rarely found elsewhere than in the blood and lymph of persons who live, or have lived, in the tropics. Thus *Demarquay* discovered it in Paris in the hydrocele fluid of a native of Havana, and *Wucherer* saw it in Bahia several times in the urine of persons with tropical chyluria.²⁹⁹ It has recently, however, been met with in temperate climates.³⁰⁰

It may infest the blood for months, or even years, without giving any sign of its presence; but commonly, by blocking or perforation of capillaries and lymphatics, it leads to haematuria and chyluria, or to haemorrhage and lymphatic exudation in different organs.

Manson discovered that mosquitoes acquire the Filaria directly from the blood of persons infested by it. Within the mosquito the parasite grows in six or seven days to the length of 1.5 mm., and can then subsist in the water where the insect lays its egg. In this way the Filaria may reach the human body through drinking water.³⁰¹ *Patrick Manson*, *Stephen Mackenzie*, and others,³⁰² have shown that in persons who suffer from the presence of *Filaria sanguinis hominis*, the parasite is to be found in the blood only at certain periods, being absent during the day, and abounding especially at night. Hence it is important that, when it is sought for, *the blood should be taken from the patient at night, and forthwith examined*.

VI. SERO-DIAGNOSIS OF THE BLOOD.—The researches of *Behring*, *Buchner*, *Gruber*, *Pfeifer*, and numerous other authors, have demonstrated the existence in human blood of peculiar protective bodies—termed alexines by *Buchner*—of still unknown nature, which are endowed with the power of killing pathogenic parasites that may effect an entry into the blood. It has also been shown that when the human organism is attacked by an infectious disease, the blood elaborates an increased quantity of the particular body capable of destroying the pathogenic organism causing that disease.

Owing to the magnitude attained by the literature of this subject, the author feels obliged to omit further literary references, and to treat of the sero-diagnosis of the blood only in so far as it has been proved to offer reliable assistance in the diagnosis of internal diseases. The credit of having been the first to definitely direct attention to the high diagnostic value of such investigations undoubtedly belongs to *Gruber*.³⁰³ Up to the present time, however, sero-diagnosis has only been utilised in

practice for a single disease—typhoid fever; and it is to *Widal*³⁰⁴ that we are indebted for arranging the existing facts in a practical clinical form—*i.e.* a form suitable for the medical man—on the basis of the observations above mentioned.

1. Details of the Widal Reaction.

(a.) **Microscopic Reaction.**—About 2–3 cc. of blood are taken from the tip of the patient's finger by pricking, and either placed in the ice cupboard, in order to recover the serum, or else separated from the latter in the centrifugal extractor. Twenty-five drops of Koch's nutrient bouillon are deposited, by means of a platinum loop, on the surface of an ordinary microscope slide, and each drop is then inoculated by the loop with a young typhoid culture, previously examined under the microscope to make sure that the bacilli are in active motion. A drop of the blood serum is added to the liquid mixture of bouillon and typhoid bacilli. The liquids are then well mixed together, and one drop is placed in the cavity of a glass slide, and examined under the microscope (oil-immersion lens).

When a typhoid affection is in question, the phenomenon termed "agglutination" by *Gruber* occurs within 3 to 15 minutes—rarely at once—the bacilli ceasing to move, and collecting together in aggregations ("clumping"); their contour disappears.

Instead of the dilution 1:25 always employed by the author, greater degrees of dilution, *e.g.* 1:100, 1:800 (*Knöspel*), 1:1000 may also be practised. A concentration of 1:10 will not, however, afford any decisive test for diagnosing typhoid, whilst the higher degrees of dilution will only give positive results in cases where the infection is grave. (See below.)

(b.) **Macroscopic Reaction.**—Five cc. of Koch's nutrient bouillon, sown with typhoid bacilli from a fresh culture—under one day old, if possible—are mixed with 5 drops of serum. In the case of blood obtained from a typhoid patient, it will be noticed that the sample turns turbid, the liquid becomes flocculent, and at the end of twenty-four hours the bottom of the test-glass will be found covered with a flaky sediment.

The literature dealing with the utility of this test is exceedingly voluminous, but the author will refrain from citing the hundreds of observations that have been made, and will merely refer to the communication from his own clinic published by *Knöspel*.³⁰⁵

2. Criticism and Diagnostic Value of the Widal Test.—In criticising the method the author will confine himself to his personal observations in more than 100 cases of typhoid, and about 100–150 cases of a non-typhoid character—diphtheria, &c.—in which this test has been applied.

In the first place, it should be mentioned that the microscopic and

macroscopic tests are of equal value. The latter, however, takes up more time, and is more expensive, a whole tubeful of bouillon being required for *each* performance of the test.

The dilution should in no case fall below 1 : 25, a rule the necessity of which is evident. Normal blood already contains a small quantity of substances capable of causing agglutination, in consequence of which the test may, and indeed *will*, give positive results with this degree of dilution.

It must, furthermore, be borne in mind that a single negative result with this test affords *no* proof that typhoid is *not* present. Consequently the blood must be tested *daily*, the reaction not occurring before the eighth day at the earliest, whilst frequently it appears only on the ninth day, and often still later.

In none of the cases (recognised by *post-mortem* examination as typhoid) referred to did the test fail to give positive indications; and, in fact, it frequently rendered a correct diagnosis possible in clinical cases which would have been otherwise very perplexing. For example, one highly febrile case was observed where the extremely purulent expectorations contained numerous mould fungi. This the author diagnosed as Pneumomycosis, but the positive results ensuing from the daily application of the Widal test indicated typhoid. Dissection revealed typhus abdominalis, particularly by the centres of purulent decay engendered by the disease in the lungs, and from which the mould fungi observed in the sputum during life were derived.

In other diseases approximating to typhus abdominalis, such as *morbus Weilii* (*Knöspel*), the reaction remained negative.

From this experience the author has no hesitation in ascribing an exceedingly high diagnostic value to the *Widal* reaction. However, the recently published observations of *Stern*,³⁰³ to the effect that the *coli* bacteria are similarly influenced by typhoid blood serum, apparently impose some limitation on this assumption. The author can confirm Stern's reports from personal experience, but maintains that these observations do not deteriorate from the diagnostic value of the Widal test. They rather indicate merely—and this is supported by other bacteriological and clinical experiences—that the form of disease termed abdominal typhus is, or under certain circumstances may be, probably a joint infection produced by the co-operative action of the typhoid bacillus and *Bacterium coli commune*.

In conclusion, it may be remarked that, in addition, a certain prognostic importance must be attributed to the Widal test, the degree of infection, and therefore, in all probability, the severity of the disease increasing in proportion with the ratio of dilution (1 : 100, 1 : 800, 1 : 1000) under which the blood continues to furnish the reaction.

VII. CHEMICAL CHANGES IN THE BLOOD.

1. Colouring Matter.⁸⁰⁷—The most important constituent of the blood is the **Oxyhaemoglobin**—the combination of the colouring matter with oxygen which is formed by the aeration of the blood in the lungs. The characteristic spectrum of dilute solutions of this body exhibits two absorption-bands between D and E (of Fraunhofer's lines). The band nearest to D is darker, narrower, and more strongly marked; that next to E is less sharply defined and broader (fig. 37).

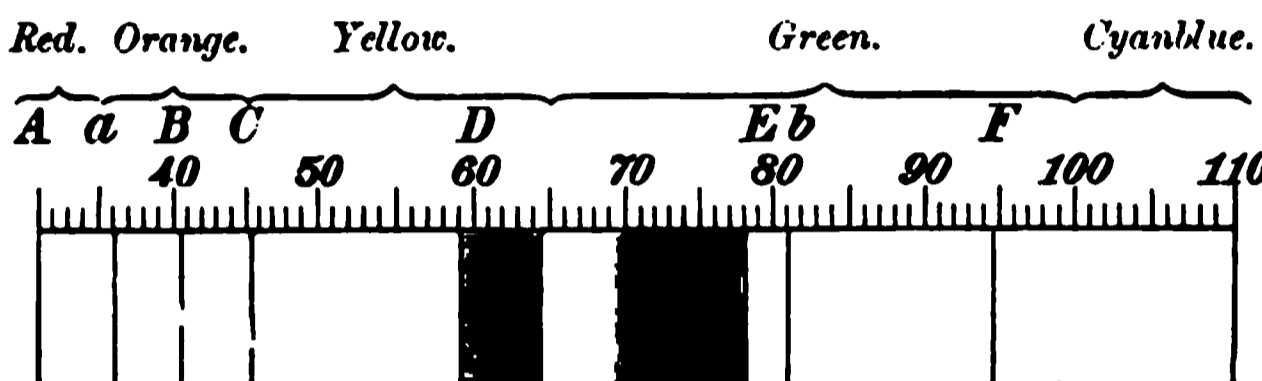


FIG. 37.—Spectrum of Oxyhaemoglobin.

When submitted to the action of deoxidising substances, oxyhaemoglobin gives place to **Reduced Hæmoglobin**. The spectrum of this body is characterised by a single band, occupying a space about midway between the former two bands (fig. 38).

Under the action of acids of all kinds, of strong alkalies, and even of CO_2 , hæmoglobin is broken up into (1) a proteid resembling *globulin*, and (2) the iron-containing body known as **Hæmatin**.

The spectrum of hæmatin in *alkaline* solution shows an absorption-

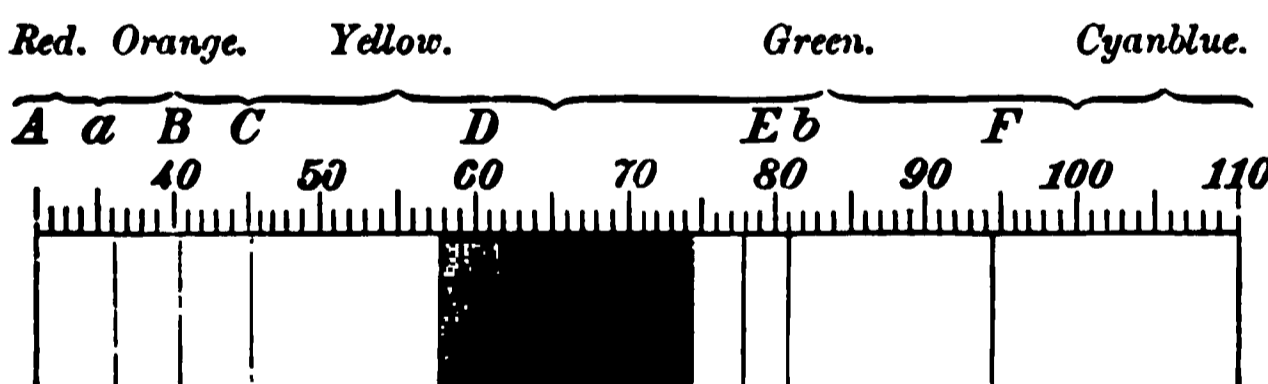


FIG. 38.—Spectrum of Reduced Hæmoglobin.

band lying between C and D of Fraunhofer's lines (fig. 39). In an *acid* solution its spectrum is identical with that of the methæmoglobin acid solution (fig. 42).

Hæmatin in alkaline solution, when treated with reducing agents, yields **Reduced Hæmatin**. The spectrum of this body (fig. 40) exhibits two absorption-bands between D and E. If the reduced solution be shaken up with air or oxygen, these bands disappear again, and the

spectrum shows once more the two absorption-bands of the alkaline solution of haematin.

Haematin in combination with hydrochloric acid forms, even from minute blood-traces, microscopic, highly characteristic, brown rhombic crystals (fig. 41). These brown rhombic crystals of haematin chloride are commonly known as haemin crystals, and were first discovered by Teichmann.³⁰⁸ They are of the utmost importance, inasmuch as their

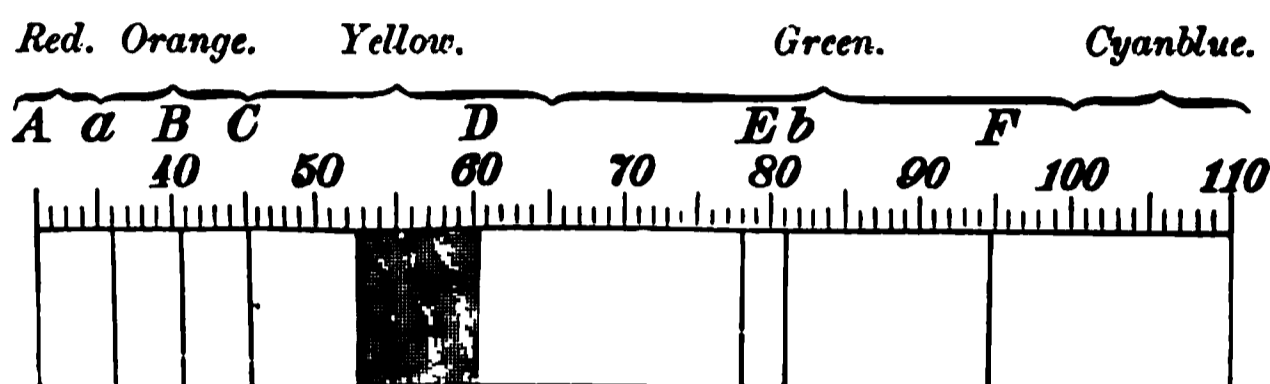


FIG. 39.—Spectrum of Haematin in an Alkaline Solution.

formation affords an admirable test for blood colouring-matter under the most varied conditions. We shall repeatedly have to revert to this later on.

To test for haemin crystals, the following plan may be adopted:—The substance supposed to contain blood colouring-matter must be dried (if not already dry), powdered, and placed upon a slide. A crystal of common salt is then added to it, and a cover-glass laid upon the pre-

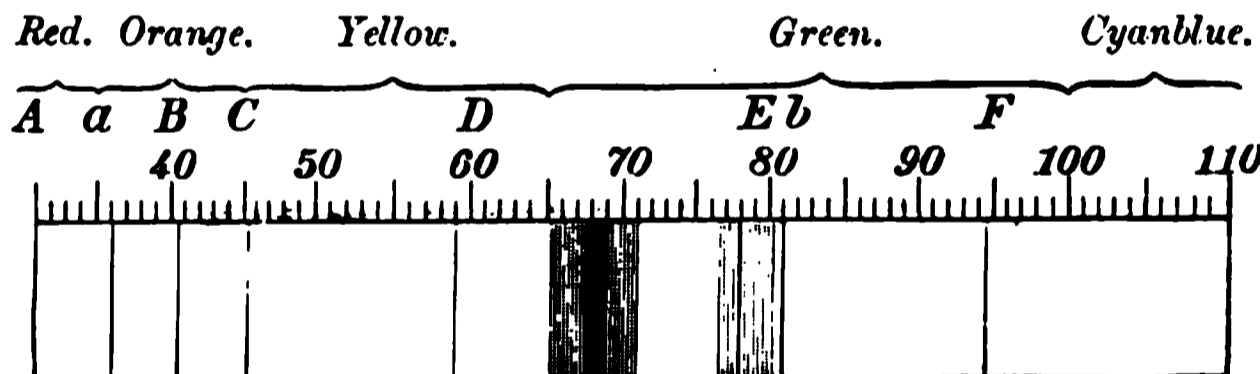


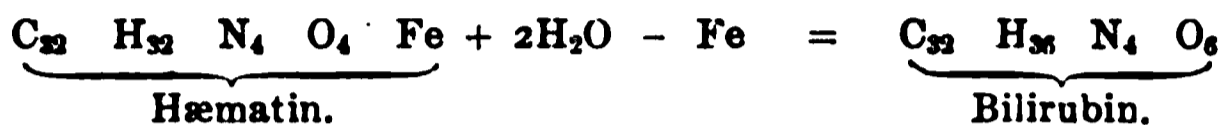
FIG. 40.—Spectrum of Reduced Haematin.

paration. A few drops of glacial acetic acid are then allowed to flow beneath the cover-glass. The whole is heated to a point below boiling, and after a little while, if the substance contains blood colouring-matter, haemin crystals (fig. 41) will be discernible by the microscope.

When an acid solution of haematin in alcohol is treated with reducing agents, a series of colouring-matter derivatives are obtained. Of these, *haematoporphyrin* (*Hoppe-Seyler*)³⁰⁹ and *hexahydro-haematoporphyrin* (*Nencki-Sieber*)³¹⁰ have been already isolated. If haematoporphyrin be acted upon with tin and hydrochloric acid in an alcoholic medium, it yields a body which cannot be distinguished chemically or by its spectrum from urobilin (*Hoppe-Seyler*).³¹¹ According to *C. le Nobel*,³¹² however, this body is otherwise not identical with urobilin.

This substance can also be obtained from bilirubin by the action of sodium amalgam (*Maly*).³¹³ There is another important derivative of haematin, which is apparently identical with bilirubin. This is **Haematoidin**, a substance first discovered by *Virchow*³¹⁴ in extravasated blood. It was afterwards found in old cerebral clots, in splenic infarctions, blood cysts, &c., and it has been met with in human urine, in the sputum, and faeces.*

From these facts, namely, that urobilin can be formed from haematin by the action of reducing agents, and that this substance can also be formed from bilirubin, *Nencki* and *Sieber* have established very simple relations between the colouring matter of the blood and that of the bile. They have constructed a new formula to express the constitution of haematin, and from this it would appear that haematin is changed into bilirubin by the addition of two molecules of water and the removal of one atom of iron, thus :—



It follows from this, according to *Nencki* and *Sieber*, that *bile pigment is formed from the colouring matter of the blood, in that its molecules lose iron and take up water.* *Latschenberger*³¹⁵ concludes, from experiments which he performed on animals, that bile pigment, or rather its antecedent, to which he gives the name of **Choleglobin**, results from the decomposition of blood-colouring matter, a dark-



FIG. 41.—Teichmann's Haemin Crystals (eye-piece III., objective 8A, Reichert).

coloured ferruginous pigment being formed at the same time. Choleglobin is elaborated both in the tissues and in the interior of cells.

It seems to the writer not unimportant to consider these views here, in anticipation of much that will have to be said later on concerning the colouring matters of the blood and of the bile, in their relations to one another.

The colouring matter of the blood forms with oxygen another compound, called **Methæmoglobin**,³¹⁶ which is distinguished from oxyhaemoglobin by the more intimate union of the O with the Hb.

The spectrum of methæmoglobin in acid and neutral solutions shows four absorption-bands (fig. 42), one well marked (between C and D), the other three in the yellow, green, and blue, being less easily seen. This spectrum, as already said, is indistinguishable from that of acid haematin in alcoholic solution. Any possibility of confounding these two bodies is, however, excluded by the fact that when *methæmoglobin* is acted upon with sulphide of ammonium, its spectrum gives place, first, to that of oxyhaemoglobin (fig. 37), and after a while to that of reduced

* See the chapters on these subjects.

haemoglobin (fig. 38); whilst, on the other hand, a solution of *haematin* treated with ammonium sulphide yields a spectrum exhibiting two absorption-bands between D and E (fig. 40). In alkaline solution the spectrum of methæmoglobin shows three bands, viz., a narrow one between C and D, but close to the latter, and two broader ones between D and E (*Jäderholm*).³¹⁷

1. Blood-changes in Dyspnoea.—All conditions which interfere with the giving off of CO₂, and the absorption of O in the lungs, are attended with certain characteristic changes in the blood.

The clinical symptoms of dyspnoea do not fall within our province. They result directly from the condition of the blood, which shows itself in the appearance of the patient. The arterial blood is laden with carbonic acid, and in consequence has a darker colour, and this imparts a bluish hue to the visible surfaces—cheeks, lips, nose, and finger-tips. Microscopical examination of the blood shows no changes of a special character. And, further, in cases of the most severe dyspnoea the blood

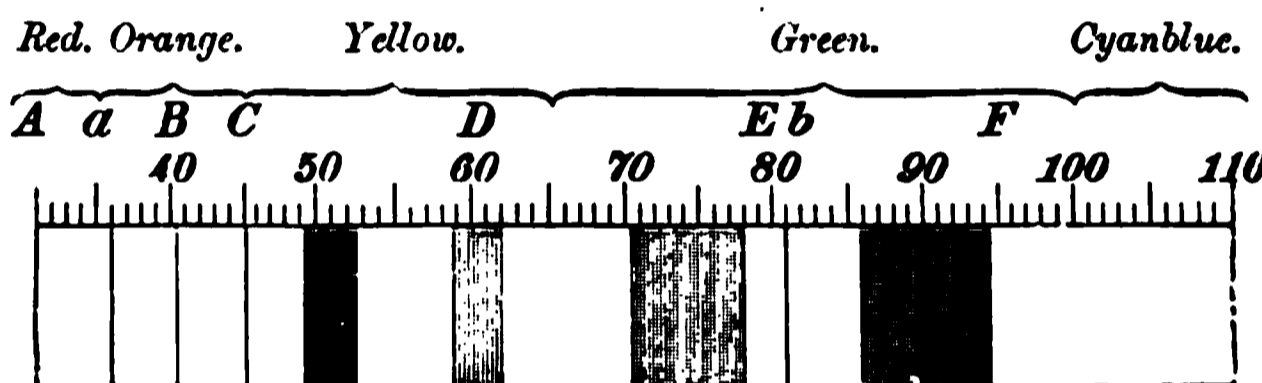


FIG. 42.—Spectrum of Methæmoglobin in Acid and Neutral Solutions.

is never so deficient in oxygen that its spectrum exhibits any considerable change, such as, for instance, the disappearance of the oxyhaemoglobin bands. *V. Loos*, by the application of Hénocque's instrument, has observed a notably diminished intensity in the oxyhaemoglobin bands in three cases of extreme cyanosis, while the proportion of contained haemoglobin was approximately normal; and it is probable that, with greater familiarity in the use of this method, we shall learn to distinguish quantitative and qualitative spectroscopic changes as a result of dyspnoea.

2. Blood-changes in Carbonic Oxide Poisoning.—In carbonic oxide poisoning the blood undergoes a change of colour which is appreciable by the naked eye. It becomes of a bright cherry-red, alike in the arteries and the veins. Spectrum-analysis shows the most important change (fig. 43). The two absorption-bands of oxyhaemoglobin are replaced by two others between D and E, but placed slightly nearer to the violet end of the spectrum. These bands indicate the union of the haemoglobin with carbonic oxide,³¹⁸ and the most important quality of this union is, that *these bands cannot, as in the case of oxyhaemoglobin,*

be made to disappear by the action of deoxidising agents (ammonium sulphide, Stokes' fluid). Carbonic oxide haemoglobin in the blood of the living subject may be recognised thus:—A few cubic centimetres of blood are taken from the patient by means of a cupping-glass, and mixed with water. Sulphide of ammonium is then added, and the solution is placed in a glass vessel with parallel sides, or, still better, the blood itself placed by means of Hénocque's apparatus before the slit of the spectroscope. If the specimen be one of blood poisoned with carbonic oxide, the two absorption-bands will remain in spite of the admixture with the reducing agent, sulphide of ammonium.

The presence of carbonic oxide in the blood can also be determined by the following chemical test:—To a quantity of blood mixed with water a 10 per cent. solution of caustic soda is added. When the mixture is slowly warmed, a cinnabar-red colour appears. Under similar circumstances a solution containing oxyhaemoglobin will turn brownish-green (*Hoppe-Seyler, Otto*).³¹⁹

A modification of this test has been suggested by *Salkowski*.³²⁰ The

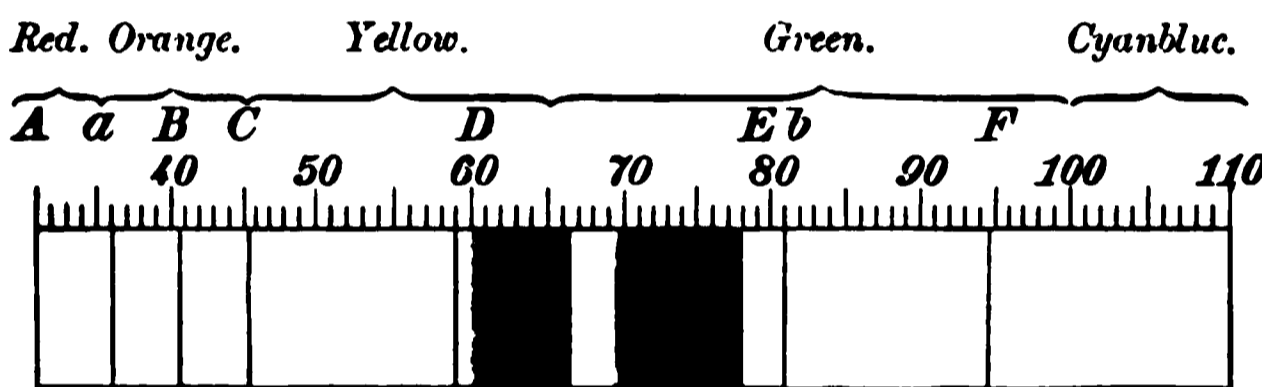


FIG. 43.—Spectrum of Carbonic Oxide Haemoglobin.

blood to be examined is diluted with water to twenty times its bulk, and a like quantity of a solution of caustic soda (sp. gr. 1.34) is added. If the blood contains carbonic oxide, the fluid turns first white and cloudy, and presently a bright red; when allowed to stand, red flakes form and settle upon its surface. In the case of normal blood, when treated in this way, a dirty brown coloration results. *Kuniyosi Katayama's* method³²¹ is to add to the blood a little yellow sulphide of ammonium and dilute acetic acid. The presence of carbonic oxide will then be shown by the appearance of a beautiful red colour, whilst normal blood so treated turns grey or greenish-grey.

Kunkel and *Welzel*³²² employ zinc chloride, or a very dilute solution of platinum chloride. With these reagents carbonic oxide blood turns a bright red, while normal blood becomes black. Other tests recommended by *Welzel* are potassium ferrocyanide, acetic acid, and tannin. *Rulmer*³²³ dilutes the blood with four to five times its bulk of acetate of lead, which causes normal blood to take a chocolate-brown colour, while if carbonic oxide be present it turns red.

3. Blood-changes in Poisoning with Sulphuretted Hydrogen (*Hydrothionæmia*).—The investigations of *Hoppe-Seyler*³²⁴ tend to the conclusion that hæmoglobin will enter into combination with H₂S, and form a substance which that author has called Sulphide of Methæmoglobin. It is, however, noteworthy that in the severest cases of poisoning with H₂S, the two absorption-bands in the spectrum of oxyhæmoglobin are never known to disappear. In such cases, the blood becomes peculiarly dark, and sometimes of a dull-green tint. And it is further remarkable that the distinction between venous and arterial blood entirely disappears (*Leirin*).³²⁵

4. Blood-changes in Prussic Acid Poisoning.—*Preyer*³²⁶ maintains that hydrocyanic acid forms a crystalline compound with hæmoglobin. However this may be, such a compound has not yet been found in the blood of men or animals poisoned with the drug. According to *Hoppe-Seyler*,³²⁷ the union of hydrocyanic acid with hæmoglobin is of a very unstable character, the resulting body readily decomposing when crystallised or in presence of putrefaction. *Kobert* describes a case of this union.

5. Blood-changes in Poisoning with Chlorate of Potash.—*Marchand*³²⁸ discovered that when potassic chlorate was taken in great quantities, the blood was profoundly altered, the most notable effect being the formation of a sepia-like decomposition product, which was afterwards shown to be identical with the methæmoglobin of *Hoppe-Seyler* mentioned above. Large doses of chlorate of potash cause the production of methæmoglobin itself in the blood, especially of children. *Stokvis* and others³²⁹ conclude from experiments upon rabbits that the exhibition of the salt is not attended with the formation of methæmoglobin in the blood of the living subject; while *Marchand* and *Cahn*³³⁰ have, in fact, obtained this result in certain animals, notably dogs. The opinion of the latter observers finds support in a clinical notice of *Lenhartz*, and also in a pathological observation recorded by *H. Hammer*.³³¹

Chlorate of potash may be easily detected by spectrum-analysis in fairly dilute solutions of hæmoglobin, and the spectrum of methæmoglobin, if present, will afford presumptive evidence of the poison. Methæmoglobin is produced also by the inhalation of nitrite of amyl (*Gamgee*) and the injection of sodium nitrite into the blood-vessels³³² (*Hoppe-Seyler*), as well as by kairin, thallin, hydrochinon, pyrocatechin, iodine, bromine, turpentine, æther, perosmic acid, permanganate of potash (*G. Hayem*), and antifebrin (*F. Müller*).³³³ [The nitrites form a compound with its oxygen, more firmly fixed than that of the oxygen in oxyhæmoglobin. They consequently tend to stop internal respiration, but are more readily displaced by the products of asphyxia than is carbonic

oxide hæmoglobin, and so again permit the aeration of the blood at the lungs.]

6. **Poisoning with Nitrobenzol.**—It has been asserted³³⁴ that in dogs poisoned with nitrobenzol the spectroscope has shown blood-changes attributable to the presence of hæmatin. It would seem, then, that in any case of suspected poisoning by this means in the human subject, the blood should be examined in this direction by the spectroscope. In a typical case of nitrobenzol poisoning which the author³³⁵ lately had occasion to investigate, the blood was of a remarkable dun colour, but presented normal microscopic and spectroscopic appearances.

7. **Hæmoglobinæmia.**³³⁶—By this term is meant the condition in which hæmoglobin is found dissolved in the blood. It is followed by *Hæmoglobinuria*, whenever the spleen and the liver are unequal to the task of eliminating the materials derived from the destruction of the red blood-corpuscles within the vessels.

The presence of free colouring matter in the blood may easily be determined thus:—A little of the blood, drawn from the patient by means of a cupping-glass, is placed immediately in a refrigerator, and allowed to remain there for twenty-four hours. If the blood is normal, perfectly clear yellowish-coloured serum will settle; whereas, if hæmoglobinæmia be established, there will be seen over the blood-clot a beautiful transparent ruby-red stratum. The spectroscope shows in the case of normal serum a feeble absorption-band in the blue part of the spectrum (at F), said to be due to lutein (*Thudichum*);³³⁷ whilst with serum containing colouring matter it shows the two absorption-bands of oxyhæmoglobin. The following method may also be employed:—Blood serum is made to coagulate by heating it to 70°–80° C. If it contains dissolved colouring matter it will appear of a more or less deep brown colour, whereas healthy blood-serum when coagulated is light yellow and of a milky turbidity. This method serves well for the detection of hæmoglobinæmia.³³⁸

8. **Recognition of Changes in the Colouring Matter of the Blood.**—The changes in the character of the blood referred to above are chiefly to be estimated by means of the spectroscope. Very perfect little instruments for clinical use have been invented by Desaga of Heidelberg, Zeiss of Jena, and Hoffman of Paris. Browning's spectroscope is also very suitable for the purpose.

To use one of these, artificial or day light is made to fall upon the slit of the instrument. The telescopic tube attached to the apparatus is focused until a spectrum is clearly defined, and if daylight be employed, the slit-like aperture is narrowed so as to bring Fraunhofer's lines clearly into view. The blood-solution to be tested is then fixed between the aperture and the light. If the fluid be too concentrated, it must be

diluted beforehand. If the light be artificial, whether from a lamp or some other source, it is well to place a little common salt or some other salt of sodium in the flame, in order to define the situation of the sodium line. Hénocque's instrument is applicable to the same purpose.

Very good results have been obtained in the investigation of blood spectra with the aid of *E. Hering's*³³⁹ "Lensless Spectroscope," an instrument which especially commends itself to the practitioner by reason of its cheapness. It has been employed by the author together with Browning's pocket-spectroscope, and shown itself to be quite as serviceable. The lensless spectroscope consists of two tubes, one sliding within the other, and of about $2\frac{1}{2}$ cm. diameter. Of these, the outer one is closed at its free end by a plate in which is a slit with a parallelogram adjustment (fig. 44 c). The two parts of the plate which support this parallel adjustment carry also a pair of clips destined to hold in position a rectangular glass vessel or test-tube containing the fluid to be examined.³⁴⁰

The tubes are lined with black, and the inner one (*a*) is provided at

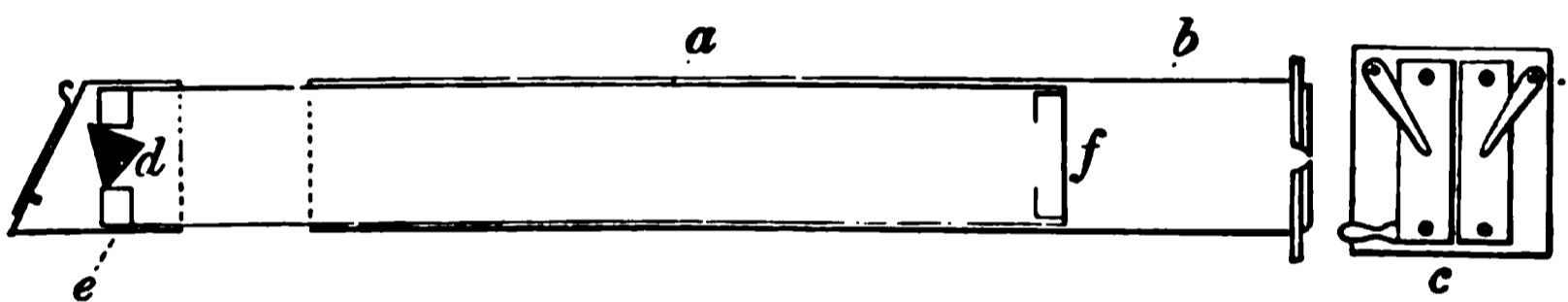


FIG. 44.—Hering's Spectroscope without Lenses.

(*f*) with a diaphragm to intercept reflected light. At that end of the inner tube which is turned towards the observer, a prism (*d*) is fixed in such a position that the spectrum is formed in a plane at right angles to the proximal end of the tube, which is oblique, not vertical, in section. In using the instrument, the eye must be directed at right angles to this section, and not in the long axis of the tube. The tubes must also be adjusted in such a manner that the spectrum is well defined and accurately rectangular.

When this is done (by manipulation of the two tubes), a small but very clear spectrum is obtained, in which the yellow is little developed, but which very plainly exhibits absorption-bands such as those of oxy-hæmoglobin and urobilin. The instrument serves admirably for the investigation of these bodies in blood and urine.*

2. Proteids of the Blood.—The proteids of the blood are diminished in all cases in which the total quantity of that fluid is greatly lessened—temporarily, therefore, in haemorrhages of all kinds.

* This instrument can be had of *Rothe*, of Leipzig, for five florins.

Since, however, the loss of blood is rapidly repaired, it is seldom that opportunity offers of recognising this condition.

It may be stated here that the quantity of blood in the body is remarkably constant, and but rarely and very temporarily exhibits any considerable variation.

A permanent diminution of proteids occurs under such conditions as disturb unfavourably the balance of waste and repair, whether of the blood itself or of its contained albumin. Thus, in all diseases which are attended with long-continued and excessive destruction of proteids in the system, these bodies are found to be proportionally wanting in the blood. It should be remarked, however, that such processes must be long continued before this effect is reached, especially where the digestive functions remain unimpaired. As a rule, diminution of proteids goes hand in hand with an unduly watery state of the blood

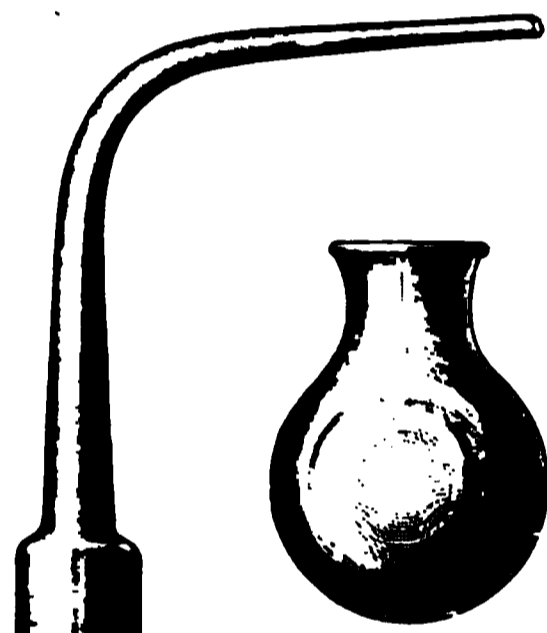


FIG. 45.—Oxidation Flask (by R. v. Jaksch).

(*hydæmia*), and is proportional to it. The author has failed to find changes in the albuminous constituent in connection with any disease of the blood, and is compelled to regard the instances recorded to the contrary as involving errors of observation.

Quantitative Estimation of Proteids.—R. v. Jaksch's Method.³⁴¹—The blood is taken by cupping-glasses, placed in a flask with a caoutchouc-stopper (fig. 45), and its nitrogen estimated by *Kjeldahl's* method (*Gunning's Mixture*). The figure expressing the content of nitrogen is multiplied by 6.25, and the result is the amount of albumin present. This method gives very accurate conclusions. It shows that, on the average, the blood of an adult contains 22.62 grms. of albumin to 100 grms. of blood. The average quantity in the blood serum is 8.86 grms. In morbid states this quantity may undergo considerable variation. As little as 8.46 grms. to 100 grms. of blood was found in a case of gastric carcinoma with severe secondary anaemia.

Results similar to these have been obtained by other methods, as by *Strauer, Stintzing, and Gumprecht*,³⁴² who used a modification of v. Bamberger's process, estimating the dried residue of the blood and deducing the proportion of albumin, and by *Maxon, Biernacki, Wendelstadt, and Bleibtreu*.³⁴³ It may be mentioned that the method employed by the latter observer is much in need of verification.

The occurrence of an absolute increase of proteids has not yet been established on satisfactory evidence. A relative increase is known to happen in such diseases as are attended with the withdrawal of water from the system more rapidly than it can be supplied, as in cholera and profuse diarrhoea. *Biernacki's*³⁴⁴ researches have shown that the withdrawal of water from the blood is by no means a constant occurrence in cholera.

In pneumonia and erysipelas there is an increase of fibrin. *Hoppe-Seyler*³⁴⁵ has devised a method for the estimation of the quantity of fibrin in the blood, which is at once simple and adapted to clinical purposes. It may be described as follows:—A beaker of about 80 cc. capacity is provided with an india-rubber cap, perforated in the middle by a close-fitting rod of whalebone. These are dried and weighed. Next, 30–40 cc. of blood, taken from the body of the patient with a cupping-glass, are placed in the beaker, which is immediately covered by the india rubber cap and its whalebone rod. The blood is then defibrinated by beating it up with the whalebone rod, allowed to cool, and weighed. The cover is removed, and the beaker filled with water and beaten up again. The fibrin is allowed to settle, washed with a solution of salt, and placed upon a filter whose weight is known. Here it is again washed with water until the fibrin is almost free from colouring matter. It is next boiled with alcohol (to dissolve fat, lecithin, and cholesterol), dried at 110°–120° C., then cooled, and weighed over sulphuric acid.³⁴⁶

*E. Ludwig*³⁴⁷ and the author have found peptones present in great quantity in the blood in leukæmia. *Devoto*,³⁴⁸ on the other hand, failed to detect peptone in this condition. In two cases investigated by him, *Matthes*³⁴⁹ determined the presence in the blood of peptone, which he further ascertained to be deutero-albumose. The author examined the blood in eight cases of leukæmia, using either Devoto's or Hoffmeister's method, and in five of these peptone was found to be present. These five were also those cases of the series which were marked by abundance of eosinophil cells and granules. The association would seem to be usual (see p. 39). A comparison of the author's investigations with that of *Matthes* establishes beyond doubt that both of them have detected the same proteid, and this, in the author's view, is peptone.

To determine the presence of peptone in the blood, it is necessary

first to remove the other proteids by the action of metallic oxides, or by coagulation with ammonium sulphate, and then to proceed in the manner indicated in the chapter on *Urine*.

3. Urea.—Urea occurs only in traces in healthy blood (*J. Picard*).⁸⁵⁰ The following method will serve to detect its presence:—Blood is diluted with 3–4 times its volume of alcohol, the mixture allowed to stand for twenty-four hours, and filtered. The precipitate is washed on the filter repeatedly with more alcohol, the filtrates are mixed, and the alcohol distilled off. The residue is treated with nitric acid, and the resulting crystalline pulp allowed to stand for some hours, when the crystal masses which have formed are pressed between folds of blotting-paper, dissolved in water, and treated with carbonate of baryta until carbonic acid ceases to form, and dried on a water-bath; the dry residue is then extracted with boiling absolute alcohol. On evaporation, the urea remains in long slender prismatic crystals belonging to the rhombic system. If enough blood has been taken (at least 200–300 cc.), or if the blood happens to contain urea in large quantities, the following tests may be performed with the resulting crystals:—

1. Dissolve some crystals in a drop of water upon a slide, add a drop or two of moderately strong pure nitric acid, and apply a cover-glass. When looked at through the microscope, the characteristic hexagonal plates of nitrate of urea will be seen.

2. To a somewhat saturated solution of the crystals add a little metallic mercury and a drop of nitric acid, and heat; gas (CO₂ and N) is rapidly evolved.

3. Heat the dried crystals in a test-tube, add a trace of caustic soda and a drop of dilute solution of sulphate of copper. A violet colour (biuret) indicates the presence of urea.

4. Over a crystal of urea pour a drop of fairly concentrated watery solution of furfural, and add immediately a drop of hydrochloric acid of 1.10 sp. gr. A play of colours takes place from yellow through green and blue to purple red (*Schiff*).⁸⁵¹

This reaction does not occur with uric acid, but is yielded by allantoin, though less promptly and clearly than by urea. It is given, moreover, by a number of other bodies.⁸⁵²

When the above method * fails to exhibit the presence of urea—as, indeed, usually happens in testing blood, on account of the very small quantity of that body which it contains—resort must be had to the more accurate process of *Hoppe-Seyler*,⁸⁵³ which can also be employed whenever a quantitative analysis of urea is attempted.

* The method has been described here because it serves well for the purpose of examining the secretions and excretions generally for urea.

[*Haycraft*³⁵⁴ recommends the following method:—20 cc. of blood are defibrinated and spread in a thin layer on a parchment-paper dialyser. This is then floated on the surface of 50 cc. of absolute alcohol in a suitable vessel, where it remains for twelve hours, the surface being kept moist by adding distilled water. To the diffusate is added an equal bulk of concentrated oxalic acid solution, and it is evaporated to dryness. To the residue is added naphtha-petroleum to remove fats. The purified residue is then dissolved in a little water and barium carbonate added, next evaporated, the residue treated with boiling alcohol and filtered. On concentrating the filtrate urea crystallises out, and may be submitted to the tests already described.]

Münzer has devised the following plan for the estimation of urea in the blood:—The latter is treated with absolute alcohol and filtered, &c., as above, the alcoholic extract evaporated, and the residue dissolved in water. This is then submitted to *Hufner's* process (see chapter on *Urine*).

The method is not, perhaps, very exact, but by its means *Münzer* has determined the presence of great quantities of nitrogenous bodies in the blood in cases of uræmia.

The author uses the following process:—The blood is weighed, placed in the flask figured on p. 81, and dried *in vacuo* at a low temperature. It is then put into an apparatus resembling Schwarz's extraction-apparatus, but provided with a well-ground cylinder to be fitted to the flask and filled with alcohol, instead of the Schwarz exhalation flask, which is all in one piece. In this the dried blood is extracted with alcohol. The apparatus as used is shown in fig. 46; A. indicating the water-supply, and B. the outflow.

The alcoholic extract is evaporated over a gentle heat *in vacuo*, and the nitrogenous matter, which consists entirely of urea, is estimated by Kjeldahl's method.

The results thus obtained are to be relied on. The following facts may be mentioned:—In a case of typhoid two analyses of the urine failed to show any nitrogen in alcoholic extract; while in pneumonia there was a trace in one instance, and none in two others. In two cases of diabetes (four analyses) 0.009–0.01 of nitrogen were found in the alcoholic extract from 100 grms. of blood.

Von Schröder's method³⁵⁵ is very delicate, but, on account of its minute details, it is hardly applicable to clinical purposes.

Urea is found in increased quantity in the blood whenever its elimination is interfered with, either by disease of the kidneys or obstruction of the urinary passages. The results already quoted will serve to show that in such conditions an excess of nitrogen may be demonstrated in the blood, and this is the expression of the urea present.

V. Schröder has shown that the formation of urea probably takes place in the liver.

4. Uric Acid and Xanthin Substances.

1. **Uric Acid.**—Garrod found uric acid to the amount of 0.025–0.175 in a thousand in the blood of persons suffering from gout. It must be observed, however, that his method of testing for this substance was far from exact.³⁵⁶

He took about 30–35 grms. of blood and allowed it to coagulate. Ten cc. of the serum were then mixed with a dilute acetic acid solution in the proportion of

1 to 10, and a delicate thread was placed in the fluid. When the blood contained not less than 0.025 per 1000 uric acid, it was found that, after twenty-four to forty-eight hours, the thread was covered with uric acid crystals.

In a few instances only did he precipitate from the blood with alcohol, and apply the murexide test. Abeles²⁵ has freed the blood from proteids by the



FIG. 47.—Schwarz's Extraction Apparatus as modified by R. v. Jakob.

Schmidt-Mulheim process, and then tested for uric acid by Ludwig and Salkowski's method. Solomon²⁶ has observed an increase of uric acid in the blood during the acute attack of gout.

For the detection of uric acid in the blood, the following procedure may be adopted.^{25a}—100-300 grms of blood are removed by cupping, and at once diluted with 3-4 times the bulk of water, heated on the

water-bath, and, when coagulation begins, treated with a few drops of acetic acid (sp. gr. 1.0335 at 15° C.) until it has a feebly acid reaction. It is left on the boiling water-bath for 15-20 minutes, then removed and filtered. The precipitate on the filter is repeatedly washed with hot water and added to the filtrate. The fluid, which should now have a slight yellow tinge, is again treated with a little acetic acid (sp. gr. as before), boiled over a flame, allowed to cool, and filtered. To the filtrate, when cold, is added a little sodic phosphate, and it is then submitted to the *Ludwig-Salkowski* process. Should it happen that the pure blood is deficient in salts, it may not coagulate on the water-bath so as to yield a sufficiently clear filtrate. This may be remedied by the addition of a little common salt.

The filtrate obtained by the *Salkowski-Ludwig* process, after the addition of hydrochloric acid, is evaporated to the bulk of 10 cc., and allowed to stand for twenty-four hours; then, if visible crystals are deposited, these are obtained on an asbestos filter, washed first with cold water and then with alcohol.³⁶⁰

1. Examine some of the crystals under the microscope. The characteristic whetstone forms, and sometimes the rhombic tables, of uric acid crystals are seen (figs. 116, 117).

2. Some of the crystals may be submitted directly to the murexide test (see below). If there should be no precipitate, or only a very slight one, after the addition of nitric acid, the fluid containing hydrochloric acid should be evaporated to dryness on a water-bath, pure nitric acid added, and this again driven off by heat. To the residue is applied, by means of pipettes, at one part a trace of ammonia, at another a little caustic soda solution. If uric acid be present a red or purple coloration develops at the spot touched by ammonia, and a blue round the soda (murexide test). Nitric acid in the test may be replaced by bromine-water, or chlorine-water, or nitrous acid (*v. Jakob*).³⁶¹ The latter serves particularly well. The use of bromine-water or chlorine-water as reagents has for its object to distinguish between uric acid and the xanthin bases.

The quantitative estimation of uric acid in the blood may be effected in the same way. The blood is first freed from albumin, and *Salkowski* and *Ludwig's* process applied. The author has frequently ascertained the presence of uric acid in proteid-free blood by means of Hopkins' process (see Chapter VII.).

The blood in health does not contain uric acid in appreciable quantity. Certain morbid states are marked by its appearance there. In croupous pneumonia it may amount to 0.008 grm. in 100 grms. of blood. It is present also in renal disease (acute and chronic nephritis and contracted kidney), in severe anaemia—finally, in all conditions which induce

dyspnoea, notably in heart-disease and pleurisy. It is absent from the blood in articular rheumatism and typhoid. It would appear that the febrile state, as such, never leads to the production of uric acid in the blood.

From what has been said it follows that the presence of uric acid in the blood is not characteristic of gout alone, and that it has not, therefore, the diagnostic significance imputed to it by *Garrod*.

2. Xanthin Bases.—Xanthin substances have been found in the blood by various observers.³⁶² They are closely allied to uric acid, and the principal are xanthin and hypoxanthin. Very probably adenin, paraxanthin, and guanin also occur. They may be detected in the filtrate after the removal of uric acid in *Salkowski* and *Ludwig's* process, by the modified murexide test already described (p. 86), and by washing the coloured residue after the application of the reagents mentioned there (*v. Jaksch*).

5. Carbohydrates.

1. Grape-Sugar.—In health the blood contains a minute quantity of sugar. To detect its presence there, the blood must first be freed from proteids, and for this purpose the old method of *Claude Bernard*³⁶³ is the best. The blood is weighed, and its own weight of crystalline sodic sulphate is added to it, and the mixture is boiled and filtered. The filtrate may be tested for sugar as below. Another method for the removal of proteids is to rub the blood in a mortar with solid ammonium sulphate, and filter. In this case also the filtrate is free from proteids. The process of *Schmidt* and *Mülheim* will also serve. *Abeles*³⁶⁴ employs for the same purpose an alcoholic solution of zinc chloride.

1. *Moore's test* will serve where sugar exists in any quantity. (See chapter on the *Urine*.)

2. *Trommer's test.* (See chapter on the *Urine*.)

3. *The phenyl-hydrazin hydrochloride test* is the best for detecting slight traces of sugar in the blood. It is conducted as follows (*v. Jaksch*):³⁶⁵—

Add together two parts of phenyl-hydrazin hydrochloride and four parts of acetate of soda; add water; and heat. Take 5 cc. of the proteid-free filtrate (which is practically a saturated saline solution), obtained by Claude Bernard's process, and, while still warm, add it to 5 cc. of the solution prepared as above. Place the mixture in a test-tube half filled with water, heat it for half-an-hour on a water-bath, and allow it to stand. Or a little of the phenyl-hydrazin salt and acetate of soda may be added in a dry state to the warm proteid-free filtrate, and the process conducted as described above. After it has cooled, when examined under the microscope, it is seen to contain separately and in clusters the

characteristic yellow crystals of phenyl-glucosazon scattered amongst colourless crystals of sulphate of soda. (See chapter on the *Urine*.)

To determine the percentage of sugar in the blood, *Fehling's* fluid may be employed (the blood having been previously freed from proteids) in the manner afterwards to be recommended for testing for sugar in the urine, and the polarimetric test* may be applied. It seldom happens, however, that the filtrate contains sufficient sugar to be appreciable with the polarimeters at present in use. *Lippich's* instrument is the most sensitive, and gives the best results in this connection.

[*Pavy's* method,³⁶⁶ if somewhat tedious, serves well for the estimation of sugar in moderately small quantity. The process may be divided into three parts :—1. 40 grms. of sodic sulphate are placed in a beaker, and 20 cc. of blood added. The beaker and its contents are weighed ; the mixture is stirred, and about 30 cc. of a hot concentrated solution of sodic sulphate added. The mixture is heated till a coagulum forms, when the fluid is poured off, the coagulum washed, and the washings added to the fluid in another vessel, which is then boiled and filtered. 2. The filtrate is boiled, and an equal quantity of the copper test solution added. The resulting suboxide of copper is collected on a glass-wool filter and washed. It is then dissolved with a little peroxide of hydrogen and nitric acid, boiled to drive off the excess of peroxide, and filtered through glass wool, which latter must be carefully washed. The filtrate contains the copper in the form of nitrate. 3. The copper solution is placed in a vessel into which a cylinder of platinum foil of known weight, connected with the negative pole of a galvanic battery, is suspended. Within this a platinum spiral is made continuous with the positive pole of the battery. The current is allowed to flow for twenty-four hours, when the cylinder is removed, washed in distilled water and alcohol, and weighed. The amount of copper deposited is the basis of a simple calculation. One part of copper corresponds to .5678 part of sugar ; hence the quantity of sugar in the blood used may be obtained by multiplying by this figure the weight of copper deposited.

*Claude Bernard's*³⁶⁷ method :—Place 20 grms. of crystallised sodic sulphate in each of six porcelain capsules, and to each add 20 grms. of the blood to be investigated. Mix the blood and salt together ; boil them till the froth above the clot becomes white, and the clot itself is free from red specks ; weigh again, and make good the loss from evaporation by addition of water. The whole is then placed in a small press, and the fluid part expressed, collected in a capsule, and afterwards filtered. The filtrate is placed in a burette. In a flask place 1 cc. of Fehling's solution, and to it add a few small pieces of caustic potash and 20 cc. of distilled water. Boil this fluid, and from the burette allow the clear filtrate of the blood to drop into the boiling dilute Fehling's solution until the latter loses every trace of its blue colour. As in all sugar estimations, the process must be repeated several times to get accurate results. Hence the reason why several capsules are prepared.

Read off the number of cc. used of the filtrate in the burette, e.g., = n cc.
The formula

$$= S \frac{8}{n} =$$

in grammes the weight of sugar per kilogramme of blood.

* See chapter on the *Urine*.

In Seeger's method,³⁶⁸ which may be taken as the type of the newer methods, the proteids are precipitated by ferric acetate. The blood is diluted with 8-10 times its volume of water, acidulated with acetic acid, and heated. When the precipitation of proteids commences, the mixture is made strongly acid by the addition of acetate of soda and perchloride of iron; then is added sufficient sodic carbonate to make the mixture faintly acid, and it is boiled, allowed to cool, and filtered through a fine cloth filter, free from starch. The filtrate ought to be clear. The residue on the filter is washed several times with water, and the remaining fluid in it expressed by means of a small hand-press. The expressed fluid is then mixed with the clear filtrate if the mixture has a slightly reddish tint from the admixture of a small quantity of blood-pigment. A drop or two of perchloride of iron is added to precipitate the last traces of the proteids. The fluid is again filtered. The sugar in the filtrate is estimated in the usual way by means of Fehling's solution.]

In diabetes large quantities of grape-sugar are found in the blood. Hoppe-Seyler³⁶⁹ describes a case in which it reached as high as 0.9 per cent. The author investigated the blood in a case of diabetes, and found 0.15 per cent. sugar by polarisation, 0.16 per cent. by titration. The researches of Freund³⁷⁰ would make it appear that a deoxidising substance—presumably sugar—exists in considerable quantity in the blood in cases of carcinoma. This has been substantially confirmed by Trinkler.³⁷¹

2. Glycogen.—Salomon and Fr. v. Frerichs³⁷² have studied the question of glycogen in the white blood-corpuscles. Gabritschewsky³⁷³ discovered that this body occurs partly in the protoplasm of leucocytes, and partly as free granules in the blood both of health and disease. For its detection the blood is spread in a thin layer between two cover-glasses, and dried in the air. A concentrated solution of gum arabic containing, in 100 grms., 1 grm. of iodine and 3 grms. of iodide of potassium, is taken, and a drop is allowed to flow between the cover-glasses. The presence of glycogen containing leucocytes, which are the same as the neutrophil cells described at p. 37, and also free granules of glycogen, is made evident by a more or less deep brown coloration, whether of leucocytes or granules. In health the blood examined after meals exhibits little or no increase of glycogen. In diabetes and leukæmia the glycogen reaction is very pronounced (see Chaps. IV., VII., VIII.). It is not, however, certain that the reaction obtained in this way with human blood is due to the presence of glycogen, and the recent observations of A. Czerny³⁷⁴ go to prove that it is derived from another substance of amyloid character existing in the blood. Huppert³⁷⁵ has devised a method based on the separation of proteids by means of a copper salt, and has thus demonstrated in a very satisfactory manner the existence of glycogen in the blood of animals. Where sufficient blood is available, this method may also be applied at the bedside.

3. Cellulose.—According to *Freund*³⁷⁶ the blood of tubercular patients contains cellulose. For the detection of sugar, cellulose, and the carbohydrates generally in the blood, the process of *Baumann* and *Udransky*³⁷⁷ may be employed with advantage. This is based upon the fact that the carbohydrates are precipitated from their watery solutions by the addition of benzoyl chloride and caustic potash, forming insoluble compounds. This combination of the carbohydrates with benzoyl chloride when treated with sulphuric acid yields furfrol, a body which may be recognised by its characteristic colour reaction.

6. Organic Acids in the Blood (*Lipacidæmia*).—Traces of volatile fatty acids are sometimes present in the blood. The author has frequently met with them. For their detection 20–30 grms. of blood are taken from the patient by means of a cupping-glass, an equal weight of sulphate of soda added, and the whole boiled and filtered. The filtrate is evaporated to dryness, and the residue extracted with absolute alcohol. The alcoholic extract in a large number of cases contained no fatty acids; but, on the other hand, these occurred in traces whenever looked for in fever and leukæmia, and occasionally in diabetes.³⁷⁸ Extraction of the blood directly with alcohol has often shown the presence of fatty acids in considerable quantity, especially in diabetes.

Lactic acid is also sometimes met with. Normal venous blood has been said to contain 0.0079 per cent. of sarcolactic acid (*Berlinerblau*). In regard to the latter and its tests, the reader may be referred to the statements of *Berlinerblau*.³⁷⁹ *L. Hougouneng*³⁸⁰ has found β oxybutyric acid in the post-mortem blood of diabetes.

7. Lipæmia.—The blood invariably contains small quantities of fat. While digestion is in progress it abounds in this substance normally; but a permanent excess of fat is also a phenomenon of certain morbid states. The blood in such cases is altered to the naked eye. It is turbid and usually paler than in health. Under the microscope a number of minute strongly-refracting globules are seen floating amongst the proper cellular elements. The white corpuscles also often contain fatty particles. [The lipæmia of diabetes gives the blood a pink or strawberry colour, and on standing a creamy layer collects on the surface.] If any doubt remains in a given case as to the nature of these particles, the addition of æther will settle the matter. If they are fatty, a drop of æther poured upon the slide will dissolve them and cause them to disappear. A method that may be adopted to the same end is the addition of 1 per cent. osmic acid solution, after staining with eosin, and subsequent dissolution of fat with æther, turpentine, toluol, or xylol.³⁸¹

Lipæmia has been met with in chronic alcoholism, chronic nephritis, and severe cases of diabetes. It also occurs in wounds of the medullary cavity of bones (embolic lipæmia) when fluid fat passes into the blood.

The author found abundance of fat in the blood of a convalescent from typhoid. His observations have led him to the conclusion that the degree of lipæmia is very variable in different diseases.

To estimate the fat in the blood he proceeds thus:—The blood is weighed, for several days extracted with æther in the modified *Schwarz* apparatus, and the æthereal extract received in a flask of known weight. The weight of the fluid after removal of the æther is that of the substances in the blood which are soluble in æther, namely, fats, lecithin, and cholesterin. In this way it was found that 100 grms. of blood contained in diabetes (three cases) 0.05–0.16, in nephritis 0.1–0.5, in typhoid 0.6, and in pneumonia 0.15 grm.

In each of the above cases the æther extract was also examined as to its content of nitrogen. In one instance only—a case of uræmia—was nitrogen found, the yield from 100 grms. of blood taken from a vein being 0.0586 grm. of nitrogen. The apparatus used was that described on p. 82.

8. Cholæmia.—By this term is meant the condition in which the constituents of the bile are found in the blood. In this connection the biliary acids and colouring matters (bilirubin) are the points of chief interest to the physician. And of these, again, the biliary acids must probably be regarded as the real toxic agents, involving great possibilities of mischief, leading to the disintegration of the red corpuscles, and, as a consequence, to haemoglobinæmia, disturbing the innervation of the heart, and slowing the pulse. As to how far cholesterin—as *Flint*³⁸² suggests—is concerned with this must be left undecided. Even where such symptoms are present, however, the quantity of biliary acids in the blood is always very small—so small at times as to escape detection by the chemical method presently to be detailed. This method, nevertheless, deserves to be described, since it will serve where the biliary acids exist in comparative abundance in the blood, and in all cases where the secretions are tested for bile. We shall therefore introduce it here:—

The blood³⁸³ to be examined must first be freed from proteids by precipitation with alcohol, or boiling it after dilution and filtering. The proteid-free filtrate is treated with acetate of lead and ammonia. The biliary acids combine with the lead and are precipitated as lead salts. The precipitate is washed with water on a filter, boiled in alcohol, and filtered. Carbonate of soda is added to decompose the lead salt. The solution is again filtered, evaporated to dryness, and the residue extracted by boiling with absolute alcohol. On evaporation, the bile salts will crystallise out, or a dull amorphous substance may remain, from which

the crystals can be derived by the addition of æther.³⁸³ The amorphous substance itself may be tested for biliary acids by *Pettenkofer's*³⁸⁴ method. This test depends upon the reaction of cholalic acid in presence of cane-sugar and sulphuric acid. To apply it, dissolve some of the crystalline or amorphous residue, obtained as above, in water; add two-thirds its bulk of sulphuric acid slowly, so that the temperature may not be raised above 60°. To the mixture now add a few drops of a solution of cane-sugar (1 to 5 of water), and a beautiful violet colour indicates the presence of biliary acids. According to *Mylius*³⁸⁵ this test depends on the formation of furfurol from grape-sugar, which then gives a play of colour with the bile acids. The reaction may also, therefore, be well displayed with furfurol.³⁸⁶

*Mackay's*³⁸⁷ physiological test may sometimes serve for the detection of these substances in the blood. It depends upon the action of bile acids, as observed in experiments on the atropinised frog's heart.

If a known quantity of blood be taken, the proportion of bile acids in the blood may be determined in the manner described above. Efforts have been made, unsuccessfully, to base an analytical test upon the polarisation phenomena of biliary acids.

Bilirubin may be recognised by taking the serum obtained from blood which has been allowed to coagulate in a sterilised glass cylinder, diluting it with water, removing the proteids by boiling, adding acetic acid, and then testing by any of the methods described later on (*vide* chapter on the *Urine*). *Hupper's* test is the best for this purpose. A large quantity of blood is required.

The same thing may be done more simply in a manner which the author has recently adopted. Blood is taken from the patients with a cupping-glass, sterilised in a fairly wide cylindrical glass, and allowed to stand for an hour or two. The serum is then drawn off with a pipette, forced through an asbestos filter by means of an aspirator, and placed in a test-tube. It is then shaken into froth. If bile pigment be present, this froth is yellow. In all other cases (as, for instance, in haemoglobinaemia, where the serum itself is tinted) the froth is quite colourless.³⁸⁸ Moreover, if more of the serum be now taken and left for three or four hours in a warm chamber at 35° C., the development of an intense green colour will mark the formation of biliverdin. A mere trace of bile pigment will cause this green colour to appear, whereas normal serum remains unchanged. A still simpler plan³⁸⁹ is to cause the blood-serum to coagulate slowly at 70°–80° C. If normal, it will show a milky yellowish tint; but if bile colouring matter be present, this is replaced by a green colour of varying intensity, according to the proportion of biliverdin formed from the bilirubin during the heating. In this way the author has demonstrated bile pigment in the blood when none could

be found in the urine. He has also observed that in nearly every case where urobilin exists in the urine, bilirubin may be found in the blood. This is an important fact, as showing that bile pigment circulating in the blood is transformed in the system—probably by the kidneys—into urobilin.³⁹⁰

The microscopical examination of jaundiced blood generally shows nothing unusual. According to *Silbermann*,³⁹¹ this condition in newly-born infants is marked by the following changes:—The red corpuscles are more or less disintegrated; they are often pale, or exhibit only round the pale centre of the corpuscle a ring of haemoglobin of normal tint. Blood-plates, macro- and micro-cytes, and poikilocytes are to be seen, together with nucleated red corpuscles and corpuscle-holding cells. In a case which came under his own observation, however, the author failed to find such appearances. They are not to be regarded as constant in jaundice.

9. Uræmia.³⁹²—When urinary products accumulate in the blood, the condition is termed *Uræmia*. The retention of these products is marked by certain well-defined phenomena, even though we are not at present able to refer them to the action of any one substance in particular. The assumption that the poisonous material is urea, or carbonate of ammonia resulting from its decomposition, has been disproved. It is now believed that the symptoms of uræmia are due in general terms to the excessive accumulation of fixed products in the blood. The interesting researches of *Bouchard*³⁹³ go far to show that they may be referred to the toxic effects of certain bodies resembling alkaloids (ptomaines) normally existing in the urine. *Stadthagen*,³⁹⁴ on the other hand, asserts that no such substances can be found in the urine. Uræmic blood shows an increased quantity of urea and extractives.³⁹⁵ In a number of cases reported by *Horbaczewski*,³⁹⁶ no increase in salts was noticed, even in the salts of potash. The author himself, as also *Peiper*, has in several instances observed that the alkalinity of the blood was greatly less than normal,³⁹⁷ and in some cases there was an excess of uric acid.³⁹⁸ There are no other characteristic changes to be noticed as occurring in the blood in uræmia.

10. Ammoniæmia.—Of this condition very little is yet known. From observations hitherto made, it would appear that the poisonous phenomena are due to the action of some body—probably an alkaloid—introduced into the system by absorption from the diseased bladder. It would be essential in such cases to examine the blood for ptomaines and toxalbumins.

11. Acetonæmia.—This term is applied to a condition in which the blood is surcharged with acetone. *Deichmüller* and the author³⁹⁹ have succeeded, by extracting the blood with æther and subsequent distillation, in separating from it a substance which gives the reactions

of acetone. In many morbid states, and especially in fevers, it is found in considerable quantity (*Reale*).⁴⁰⁰

12. Changes in the Inorganic Elements of the Blood.

1. **Inorganic Salts.**—The blood contains about one-half per cent. of chloride of sodium,⁴⁰¹ and this quantity remains constant, whether much salt be taken with the food or not. Moreover, *Schenk*⁴⁰² has shown that in fevers, as, e.g., pneumonia, where the chlorides disappear from the urine, the proportion of salt in the blood is not notably altered.

In rickets and osteomalacia the salts are diminished.

The blood of tubercular patients, according to *Freund*,⁴⁰³ is relatively deficient in sodium salts and phosphates, whilst at the same time the salts of potash are increased.

The tests and methods for the qualitative and quantitative analysis of the salts of the blood are to be found in the various text-books of physiology and physiological chemistry.⁴⁰⁴

2. **Watery Constituent of the Blood.**—The blood of an adult contains on an average 77.33 per cent. of water. The quantity is increased in anaemia, and is in an inverse ratio to the amount of proteid present. Thus, in the case recorded at p. 81, where the proportion of albumin was the smallest noted (8.46 per cent.), the quantity of water was also the greatest, viz., 90.01 per cent. To estimate the amount of water, the blood is weighed, and dried at 110° C. until it ceases to lose in weight. This method is not entirely accurate, but is nevertheless better than *Stintzing's*.⁴⁰⁵

[According to *Hutchinson*,⁴⁰⁶ the alkalinity of the blood is in apparent relation to its watery constituent, and this he refers to the greater ease with which watery blood neutralises the acid in the usual, and especially in the percolation tests. He accounts in this way for the seemingly increased alkalinity of the blood in anaemia when it is estimated by *Haycraft* and *Williamson's* method.]

CHAPTER II

THE BUCCAL SECRETION

THE saliva is a mixed secretion, derived in part from the mucous glands within the mouth, and partly from the parotid, submaxillary, and sublingual glands, which open by ducts within that cavity. Any disproportionate activity, whether in health or disease, of one or other of these glands will be attended with a corresponding modification of the physical and chemical characters of the saliva.¹

[To obtain the saliva fairly pure, the patient should be made to wash his mouth with a warm solution of bicarbonate of soda, and afterwards with cold water. The inside of the mouth should then be lightly touched with a glass rod, moistened with dilute acid, and the secretion collected.]

I. NAKED-EYE APPEARANCES OF THE SALIVA.—The saliva, when freshly taken from the mouth, is a colourless or light blue fluid, usually somewhat thick and stringy. When allowed to stand for some time, it settles into two layers, of which the lower one is quite cloudy and turbid, and contains in the greatest abundance the morphological constituents presently to be described.

The reaction is distinctly alkaline. [The amount secreted daily is variously stated at from 800 to 1500 grms.]

II. MICROSCOPICAL APPEARANCES.—The saliva, when examined with the microscope, is seen to contain certain morphological elements in varying proportions. These are:—

1. Salivary Corpuscles.—These bodies resemble white blood-corpuscles, but are larger, and their protoplasm is usually very granular.

2. Red Blood-Corpuscles.—These are seldom met with, and when they occur are readily recognisable.

3. Epithelium.—Usually in the form of large irregular squamous cells, derived from the mucous membrane of the mouth and tongue. The quantity of epithelium to be found in the saliva varies greatly in health; and the cells exhibit much difference in shape, according as

they come from the superficial or the deeper layers of the mucous membrane. They are, however, easily known by their polygonal shape and relatively large size.

4. Fungi.—Mould- and yeast-fungi are very seldom seen in the saliva in health ; when they occur, it is as an accidental constituent, probably introduced with the food. In disease, however, their presence is frequent. Fission-fungi, on the other hand, are met with in great number and variety in healthy saliva. There are to be seen, thickly scattered through the secretion, smaller or larger colonies of micrococci, of which some possess the property of staining reddish in a solution of iodine and iodide of potassium. *W. D. Miller*² describes four varieties of these, which he has named *Bacillus maximus buccalis*, and *Iodococcus magnus*, *parvus*, and *vaginatus*. Other bacilli, of varying size, take a bluish-red

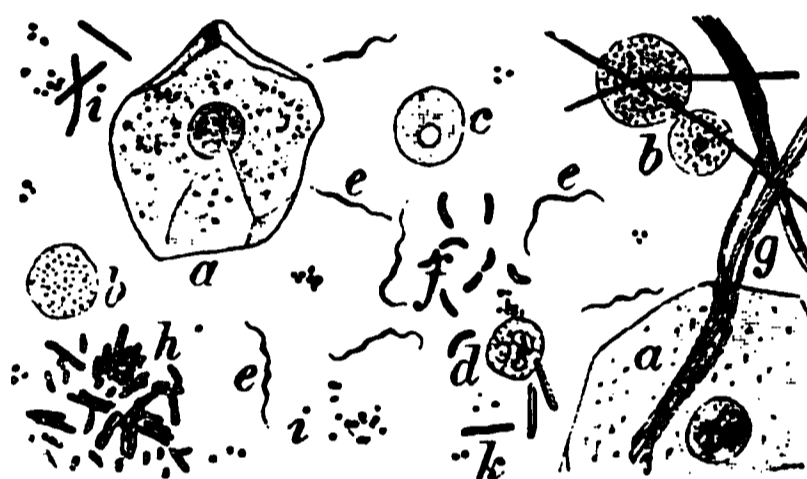


FIG. 47.—Buccal Secretion, prepared by Friedländer's and Günther's methods (eye-piece III., objective Reichert $\frac{1}{4}$; homogeneous immersion; Abbe's mirror; open condenser).

- | | |
|--|--|
| a. Epithelial cells.
b. Salivary corpuscles.
c. Fat drops.
d. Leucocytes. | e. Spirochete buccalis.
f. Comma bacilli of the oral cavity.
g. Leptothrix buccalis.
h, i, k. Different forms of fungi. |
|--|--|

colour with the same reagent. There is also an organism called the Spirochæte buccalis, which occurs in extremely mobile spiral threads, very closely resembling the spirillum of relapsing fever ; from this it is distinguished chiefly by its greater breadth and by the smaller number of its coils. Forms resembling the comma bacillus are frequently found in the saliva (*Levis, Miller*).³ They have been obtained in considerable numbers from the secretion (*Vignal*).⁴ As many as twenty-one different micro-organisms have been separated by the ordinary methods (see Chapter X.), and cultivated on plates and by inoculation, and their behaviour in various food media observed. *Biondi*⁵ has recently been engaged in such research. According to *W. D. Miller*,⁶ the following list is a summary of the pathogenic fungi which have hitherto been found in the buccal cavity, part of these having also been isolated by cultivation methods :—*Leptothrix buccalis*, *Vibrio buccalis*, *Spirochæte*

dentium, *Micrococcus tetragenus*, *Micrococcus de la rage* (*Pasteur*), *Micrococcus* of septicæmic sputum, the fungus designated 8 by Miller, the bacillus of decaying teeth, *Bacillus crassus* sputigenus, *Bacillus salivarius* *septicus*, two pathogenic fission-fungi not yet cultivated (*Kreibohm*), *Staphylococcus pyogenes*, *albus* and *aureus*, and *salivarius* *pyogenes*, *Coccus salivarius* *septicus*, and *Bacillus septicus* sputigenus.

*Miller*⁷ has cultivated over fifty different fungi obtained from the mouth. A special interest attaches to the presence in the mouth of healthy persons of the bacillus of sputum-septicæmia. Pure cultivations of this bacillus have been made by *Klein*, *Miller*,⁸ and *Fränkel*, and it would appear to be the same which the researches of *Fränkel* and *Weichselbaum* indicate as the specific cause of pneumonia (see p. 131). The observations of *Löffler*, *Vetter*, *E. Doernberger*, and *Welch*⁹ have shown that other formidable parasites, as the bacillus of diphtheria, *Staphylococcus*, and *Streptococcus pyogenes*, are normally innocuous denizens of the mouth.

To show Spirochæte buccalis, a drop of pure saliva should be examined as it is with a good oil-immersion lens, an Abbe's condenser, and a narrow diaphragm. It may be seen also in a preparation stained by *Günther's* process.

Under pathological conditions other pathogenic fungi are to be found in affections of the mouth; as, for instance, thrush-fungus, *Actinomyces*, and the bacilli of tubercle. *Fränkel*¹⁰ has obtained the bacillus of typhoid from the lingual glands in a case of death from that disease, and doubtless our acquaintance with such forms will extend with our knowledge. *Trumpp*¹¹ found diphtheria-bacilli among healthy persons both in the mouth and elsewhere.

III. CHEMICAL CONSTITUTION OF THE BUCCAL SECRETION.

—This varies even in health with the activity of the different glands by which the fluids are secreted. There are to be found traces of albumin coagulable with heat, mucin, and occasionally sulphocyanide of potassium (CNSK). The saliva contains, further, a ferment which changes starch into sugar, and a trifling amount of salts. Oxygen, nitrogen, and carbonic acid gases have been obtained from parotid saliva (*Külz*).¹²

It is seldom that opportunity offers for a chemical examination of the saliva in disease. The quantity is then generally diminished rather than increased; and further, it is very difficult to obtain a pure secretion from the patient.

Ptyalism (*see below*) is the only morbid state in which a large quantity of pure saliva can be had. To examine the saliva, the patient should be made to rinse the mouth with water carefully after each meal, and the secretion collected for twenty-four hours in a clean vessel. The reaction

may be tested with litmus paper and the specific gravity taken. It will be found to be alkaline, and of sp. gr. 1.002-1.006. A portion of the fluid may next be tested for albumin in the manner to be described in the chapter on *Urine*.

Another portion may be tested with solution of ferric chloride for sulphocyanides. Should such be present, a bright-red colour appears, which does not disappear either with heat or on the addition of acid. If the red colour is not thus obtained, 100 cc. of the saliva should be concentrated on a water-bath and tested as before. [Meconic acid yields a similar cherry-red colour with ferric chloride, and this may be obtained from the saliva in cases of opium poisoning. The addition of mercuric chloride causes the colour due to sulphocyanide to disappear, whilst that from meconic acid is unaffected by it.] *Colosanti*¹³ advocates the following method:—The saliva is precipitated with alcohol and filtered. The filtrate is evaporated on the water-bath, and the residue dissolved in water. Cupric sulphate is then added. If sulphocyanide be present an emerald-green colour develops.¹⁴

Sugar may be sought for in the manner recommended for its detection in blood, No. 3 (*v. supra*, p. 87).

The presence of diastatic ferment may be shown thus:—5 cc. of saliva are mixed with 50 cc. of starch solution, and placed in a warm chamber or in a water-bath heated to 40° C. When examined after one hour, the fluid (which of course must have been tested beforehand to ascertain the absence of sugar) will give all the reactions of grape-sugar if amylolytic ferment be present.¹⁵

Nitrites often occur in saliva. They may be detected by adding to a little of the fluid a mixture of starch paste, iodide of potassium, and dilute sulphuric acid, when, if nitrites be present, an intense blue colour will be seen. A very useful test for nitrites has been suggested by *Griess*.¹⁶ To a specimen of saliva diluted with five times its bulk of water, a few drops of sulphuric acid are added, and then metadiamido-benzol, which melts at 63° C. The appearance of an intense yellow colour shows the presence of nitrites.

As a further test, *Griess*¹⁷ acidifies the fluid to be tested with sulphuric acid, and adds a small quantity of a solution of sulphanilic acid, as also a few drops of a solution of sulphate of naphthylamin coloured by coal-tar. The presence of nitrites is indicated by a red colour.

Another test for nitrites is that of *Nosvay*,¹⁸ recommended by *Lunge*:—0.5 grm. of sulphanilic acid is added to 150 cc. of dilute acetic acid, and this is mixed with a solution of naphthylamin (1 grm. of solid naphthylamin in 20 cc. of boiling water). The fluid is poured off from the blue precipitate, 150 cc. of dilute acetic acid added, the two well mixed, and

the mixture preserved in an air-tight bottle. The fluid to be tested for nitrites is treated with this reagent and heated to 80° C. The result should be a red coloration.

IV. CONSTITUTION OF MORBID SALIVA IN GENERAL.—

The quantity of saliva is diminished during inflammation of the salivary glands in febrile disorders and diabetes, and often also in nephritis. [In high fever no saliva is secreted. That of moderate fever is thick and scanty, and usually acid, and with the rise of temperature its diastatic action is lessened. The secretion is arrested by certain drugs, notably by belladonna.] It is increased in inflammation of the mouth, by the action of certain poisons—as e.g., pilocarpin and mercury¹⁹—[in trigeminal neuralgia] and sometimes by the irritation of carious teeth. The excessive secretion which attends poisoning by acids and alkalies is rather due to irritation of the ducts than to any specific action on the salivary glands. A long-continued flow of saliva will sometimes occur without its being possible to ascribe it to any of the causes mentioned. In such cases probably the disturbance is due to some obscure changes in the innervation of the glands. Salivation has occasionally been recorded as occurring in pregnancy (*Schramm*).²⁰ [It is frequently met with in hysterical women.]

These are the cases (referred to above) which afford a favourable opportunity for chemical analysis of the saliva.

In a case of ptyalism which the author observed, analysis of the saliva showed that it contained 995.2 grms. of water and 4.8 grms. of solids. Its reaction was alkaline. It held a small quantity of *mucin*, traces of serum-albumin, and some sulphocyanides. The iodide of starch test showed the absence of nitrites; and no sugar was detected by phenyl-hydrazin or other reagents (*Salkowski*).²¹

Certain diseases are attended with notable qualitative changes in the saliva. [Its reaction may be acid in diabetes, acute rheumatism, and mercurial poisoning.]²² In nephritis considerable quantities of urea have been found in it by *Wright*, *Picard*, *Rabuteau*,²³ and *Fleischer*.²⁴ For its detection Fleischer employs the following method:—An alcoholic extract of the saliva is made and filtered, the filtrate evaporated, and the residue dissolved in amylic-alcohol. Crystals of urea remain after evaporation, and may be recognised by any of the tests described at p. 83.

*Boucheron*²⁵ found uric acid in the saliva of uræmic patients by employing the murexide test (p. 86). In such cases the author has tested the saliva in the way described at p. 86, having previously promoted its secretion by administering pilocarpin, and has never succeeded in finding uric acid there.

Bile pigment and sugar have not yet been found in saliva. Even in that of diabetic patients there seems to be no sugar. In three cases of diabetes the

author has carefully tested the secretion after the injection of pilocarpin, by means of the phenyl-hydrazin test, but in each case without any result.

Certain drugs, and amongst them iodide and bromide of potassium, are readily detected in the saliva soon after they have been taken into the system.²⁵ (See chapter on Urine for the method of investigating this body.)

[The following method for the detection of mercury in the saliva is adopted by Ralfe.²⁶ To the saliva secreted during twenty-four hours dilute HCl is added. The mixture is heated for two hours in a water-bath, filtered, and the filtrate concentrated to half its bulk. The precipitate on the filter is placed in a beaker three parts full of dilute HCl and heated, while small quantities of potassium chlorate are added, and the mixture stirred to dissolve organic residue. It is then filtered and the filtrate added to the previous one. The fluid is concentrated to one-fourth its bulk. It contains all the mercury as bischloride. To test for this.—1. Place a drop on a gold or copper coin and touch this with the blade of a knife; a bright silvery stain results. 2. Boil some of the fluid with pure copper foil; mercury is deposited on the foil, and may be volatilised in a test-tube.]

[**V. THE SULPHOCYANIDE OF THE SALIVA.**—The origin and purpose of this salt in the economy have long been a subject of speculation to physiologists. The researches of Dr. S. Fenwick²⁸ have invested the matter with a new interest, and their results go to prove that we have in the variations of its quantity a valuable index to certain states of the system. He has collected the records of a large number of cases in which the saliva was examined, and the quantity of sulphocyanide carefully compared. This was done by noting the colour produced by adding to the secretion a certain quantity of a standard solution of perchloride of iron. For the purposes of comparison, the tint so obtained with a mixture of the secretions from many healthy persons is taken to indicate the normal amount of the salt, and a scale of colours is prepared by evaporating and diluting the fluid to certain proportions, and copying the tints in each case.

As a result of his observations, Dr. Fenwick concludes that the sulphocyanide of the saliva is a measure of the functional activity of the nutritive organs, and that it is increased in general whenever an unusual demand is made on them by the necessities of the system, provided those organs are capable of responding to the call,—in the early stages of acute inflammation, of cancer and phthisis, in acute congestion of the liver from alcohol and over-feeding, in acute rheumatism, gout, and urticaria, and in convalescence from typhoid and similar diseases. The quantity is diminished in all conditions where the nutritive organs are unable to supply the requirements of the system, in the later stages of phthisis and malignant disease, in long-continued diarrhoea and dysentery, in jaundice from obstruction, in lead poisoning, and in ascites and similar conditions impeding the portal circulation, and where the assimilation of food is imperfectly performed. Where, in connection with articular

rheumatism, the sulphocyanide is greatly in excess, a tedious recovery is to be expected, and frequent relapses may be feared.

Dr. Fenwick believes that the sulphocyanide is derived from the decomposition of biliary compounds (? taurocholate of soda).

A more accurate method of ascertaining its amount is to collect the saliva secreted during five minutes, add the tincture of perchloride of iron in the proportion of one drop to a drachm, and filter. The colour obtained in this way may then be compared with solutions of sulphocyanide of iron carefully graduated. The bottles containing the tests, and the solutions to be compared with them, should be of exactly equal size. The flat-sided vessels or haematinometers used for the spectroscopic examination of blood will serve well.]

VI. THE SALIVA IN CERTAIN DISEASES.

1. Catarrhal Stomatitis.—This affection is regularly attended with a much-increased flow of saliva, which when examined microscopically is found to contain an excess of epithelium and many leucocytes, but is otherwise unaltered.²⁹ Its reaction is generally, but not always, acid.

2. Ulcerative Stomatitis.—(*Mercurial, Scorbutic, &c.*).—The saliva is foetid, dark brown in colour, and strongly alkaline. It is loaded with tissue débris, leucocytes, broken-down red blood-corpuscles, and various forms of fungi in abundance. *Fröhwald*³⁰ believes that a specific bacillus which he discovered has a special relation to ulcerative stomatitis. His contention, however, remains to be proved.

3. Thrush.—The presence of this fungus in the mouth demands a more detailed notice.³¹ It occurs most frequently in children, but is common also in adults, especially in association with tuberculosis. *Freudenberg*³² has detected it in healthy persons. It used to be taught that the saliva of thrush is always acid; but it is still a matter of doubt whether the acidity is not due rather to the presence of other micro-organisms than to the action of the thrush fungus. *Kehrer* has shown that the latter parasite will thrive well in a medium where no free acid exists, as in lactate of sodium or potassium. The outset of the disease is marked by the formation of white patches on the mucous membrane, and when examined microscopically these patches are seen to enclose sharp-bordered oval cells, each having one or two nuclei. The cells are disposed in groups of two or three. After the lapse of some days the patches run together, and form a membrane which may cover the entire mucous surface of the mouth, and even line the fauces and œsophagus.

The membranes are at first firmly adherent, but later on loosen, and may then be easily detached. When examined microscopically they are

seen to consist of epithelial cells, leucocytes, and débris, amongst which the parasite appears as branching ribbon-like forms composed of long segments. Each segment usually contains two strongly refractive nuclei embedded in a clear substance, one at either end. The segments vary in length, and grow shorter towards the extremities of the parasite. They are for the most part homogeneous, but occasionally finely granular. There are also to be seen the oval bodies figured below, which are thought to be the spores (gonidia) of the fungus.

There is still much dispute as to the place of the thrush fungus in the vegetable kingdom. *Rees*²³ refers it to the yeast fungi; *Grawitz*²⁴ supposes it to be identical with the fungus studied by Cienkowsky; and *Plaut*²⁵ opposes this view, regarding it (with both the above-named authors, and also *Boginsky*²⁶ and *Klemperer*,²⁷ as a yeast fungus.²⁸ According to the more recent investigations of *Plaut*²⁹ the thrush fungus is identical with the widely distributed *Monilia candida*.

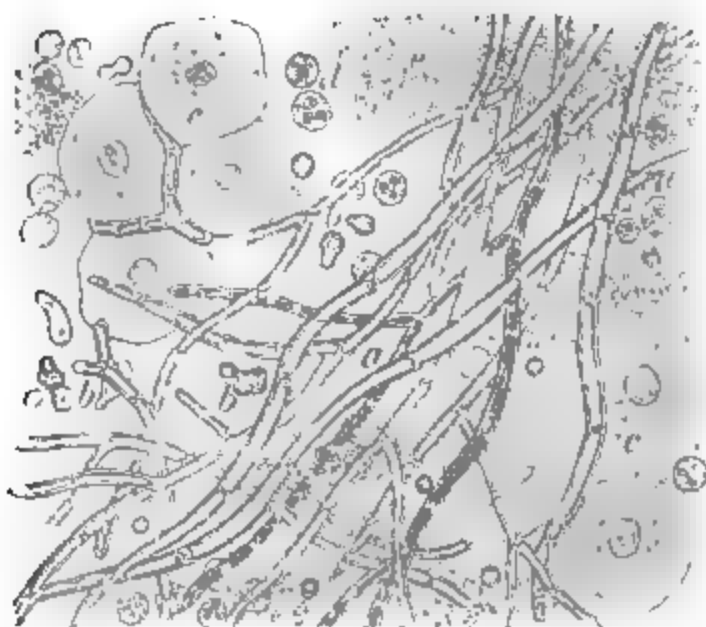


FIG. 48.—a. Thrush fungus; b. Gonidia; c. Epithelial cells; d. Leucocytes; e. Debris. (From the mouth of a patient with a weak heart; eye-pleco III, objective 8A, Reichert).

The observations of *Langerhans*,⁴⁰ and more recently of *Charrin* and *Ostrowsky*,⁴¹ give reason for the belief that this parasite, though generally innocuous, may also in the human subject be an agent in suppuration.

The fungus can be easily examined by placing part of the loose membrane with a little glycerine under the microscope.

When pus containing *Actinomyces* has been discharged into the mouth, the micro-organisms can be found in the saliva. For its recognition, see the chapter on *Pus*.

Fischer and *Hauser*⁴² have repeatedly found sarcines in the buccal mucus of wasting diseases.

VII. DEPOSIT ON THE TEETH.—If a little of the tartar be removed from the teeth with a spatula and examined, it will be seen to abound mainly in micro-organisms. These comprise:—

1. *Spirochete buccalis* (mentioned above), in small numbers.
2. *Leptothrix buccalis*. Long bacilli, usually segmented and arranged in large ribbon-like bundles. They stain bluish-red in iodine-potassic-iodide solution (fig. 49). This micro-organism has been named *Bacillus maximus buccalis* by *Müller*.⁴³ He rejects the view that it has the power of penetrating dentine, but holds with other observers⁴⁴ that dental caries is brought about by various fungi, both cocci and bacilli, which generate acids and destroy decalcified dentine. Mixed up with the masses of *Leptothrix* are usually to be seen shorter bacilli, which do not stain in the iodine solution.

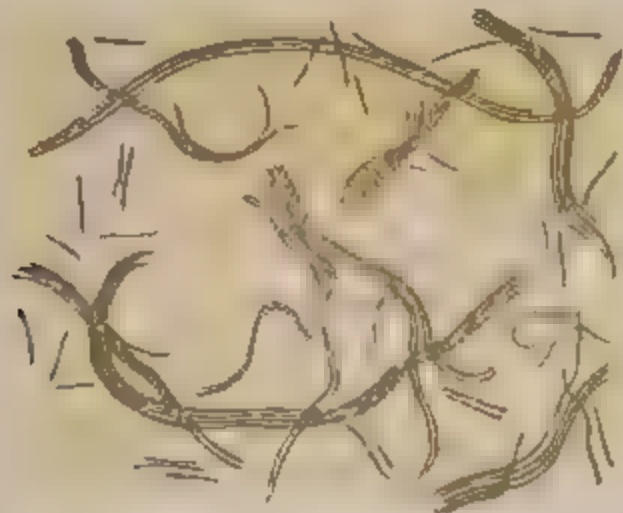


FIG. 49.—*Leptothrix buccalis*.
The preparation was stained with iodo-potassio-iodide solution (eye-piece III., objective 8A
Reichert.)

3. Various forms of micrococci, occurring both separately and in colonies.
4. A large number of leucocytes and epithelial cells, usually showing advanced fatty degeneration.

VIII. COATING OF THE TONGUE.—1. In severe infectious diseases the tongue is coated with a brownish fur, which consists partly of the remains of food and partly of dried blood. Microscopical examination of the coating removed from the tongue exhibits, in such a case, a profusion of epithelial cells and hosts of fungi of various forms. In addition to these, there is a multitude of dark cellular bodies, derived doubtless from the corneous and exfoliated epithelium of the part (*Bizzozera*).

Schuch,⁴⁵ again, has called attention to the occurrence of a black fur, which is probably conditioned by the formation of pigmented papillæ on

the tongue.⁴⁶ *Ciąglinski* and *Hewelke*⁴⁷ traced a similar appearance to a pigment-forming yeast fungus.

2. The tongue of infants is normally coated with a white fur, and a similar appearance is found in adults when the stomach is deranged. Microscopical examination shows epithelium, a few salivary corpuscles, and very many fungi.

[Dr. Dickinson⁴⁸ has recently investigated the nature and significance of the various morbid coatings of the tongue. He believes that a just conclusion cannot be arrived at from the inspection of material scraped from the surface, and his method was to obtain *post-mortem* sections through the substance of the tongue, associating the microscopical with the naked-eye appearances during life. Thus it is seen that the different varieties of coating distinguished as "stippled," "coated," "plastered," "furred," and "encrusted," are all alike derived from excess and alteration of the epithelial elements of the tongue; the change, where most profound, extending first between the papillæ, and then deeper, with hypernucleation of the deep cells of the corium and the diapedesis of leucocytes. The presence of non-pathogenic fungi is common to all, and may be regarded as accidental; even the thrush fungus occurs independently of the grosser changes to which it commonly gives rise. The conditions of dryness and moisture greatly modify both the character of the coat and its significance as a symptom. The colour of the encrusted brown variety is due to dryness alone, while at the same time it must be mentioned that a profusion of micro-organisms is especially associated with this form. Clinically the fact of most importance is the thickness and exuberance of the coating, and a comparison of instances has shown that such redundancy is especially connected with pyrexia.]

IX. COATING OF THE TONSILS.—The examination of morbid deposits upon the tonsils is sometimes of the utmost importance in diagnosis. Such deposits may be due to the action of chemical irritants, as ammonia vapour, acids and alkalies, or to the various micro-organisms, Streptococci, Staphylococci, and the diphtheria-bacillus.

1. **Deposits due to Streptococci, Staphylococci, and Diphtheria-bacillus.**—It is impossible, at least at the outset and in the case of adults, to determine by simple inspection alone whether a deposit is the result of croupous tonsillitis or of true diphtheria. For this purpose a bacteriological examination must be made. The points to be decided are these:—

(a) Whether there are present only Streptococci, Staphylococci, and cocci.

(b) Whether these micro-organisms are present together with diphtheria-bacilli.

(c) Whether the membrane or deposit contains diphtheria-bacilli alone or almost alone.

Staining and cultivation methods will show the presence of micro-organisms in the coating of the fauces, whatever its origin and import; only in benign cases the micro-organisms are those which normally

inhabit the mouth, and are innocuous, generally speaking, or, at most, include certain others, as, for instance, the micrococcus of mouse septicæmia, which are pathogenic in animals.

The membrane covering the tonsils both in croup and diphtheria is composed of glistening homogeneous fibrin, disposed in the form of a network, of which the meshes vary in shape and size, and enclose epithelial cells, blood- and pus-corpuscles, and micro-organisms of every description. The differences between croupous and diphtheritic membrane cannot be distinguished by microscopic examination alone, as was formerly taught. In both cases whitish layers are found on the tonsils. It should be mentioned, however, that *E. Wagner* discriminates between croupous and diphtheritic tonsillitis, in that the removal of the membrane in the former affection leaves the underlying tissues simply hyperæmic and infiltrated with serum, while in the diphtheritic form a haemorrhagic or even sero-purulent infiltration remains.

The observations of *Roux* and *Yersin*, *Zarniko*, *Spronck*, *Wintgens*, and *van den Brink*, *Paltauf* and *Kolisko*, *Escherich*, *Klein*, and *Beck*,⁴⁹ fully confirm the view that the bacillus first discovered and described by *Klebs* and *Löffler*⁵⁰ is to be regarded as the cause of diphtheria.⁵¹

*G. v. Hoffmann*⁵² has shown that a micro-organism, morphologically and in its life-history closely resembling the bacillus of *Klebs* and *Löffler*, occurs in the mucous membrane of diphtheria patients. This he has named the pseudo-diphtheria-bacillus. *Kolisko* and *Paltauf*, whose observations have been confirmed by those of others, have again called attention to a multiple infection in this disease—Strepto- and Botryococci being found in the interstices of the tissues, and diphtheria-bacilli on the surface. These discoveries tend somewhat to discount the significance of the diphtheria-bacillus as distinguished by Löffler's indications. They have, however, a very great diagnostic importance, and in connection with Behring's serum remedy bear indirectly upon the therapeutics of diphtheria. *Escherich*⁵³ maintains that even in diphtheria the pseudo-bacillus appears less frequently than Hoffmann supposed, and that its presence is inconstant. It has likewise been shown, in a highly interesting series of observations by *Westerbrook*, *Wilson*, *Daniel*, and *Adair*,⁵⁴ that similar atypical diphtheria-bacilli also appear in children not suffering from diphtheria.

More hopeful are the researches of *Roux* and *Yersin*, *Brieger* and *Prankel*, *Wissermann*, *Martin*, and *Proskauer*,⁵⁵ which teach us that these fungi elaborate proteid substances (toxalbumins) of a very poisonous nature. When a method has been discovered by which it will be possible to isolate these substances, either from pure cultivations of the fungi or from the diseased tissues, we shall probably be in possession of a more certain and simpler, and for that reason clinically more serviceable, diagnostic resource than is afforded by the present cultivation processes.

[*Sidney Martin*⁵⁶ has obtained from among the products of the bacillus of diphtheria definite substances, which he believes to be separately the agents in the development of different symptoms. These substances are mainly two albumoses and an organic acid. The albumoses he obtained from the diphtheritic membrane, from pure cultivation of the bacillus, and from the spleen and blood of persons suffering from the disease. The injection of these albumoses upon animals was followed by febrile disturbance, and later by degeneration of peripheral nerves, of the heart muscle and of other organs. The organic acid in question has similar but less actively poisonous properties. Martin's theory may be stated thus: the bacillus fastens upon a fibrinous exudation in the throat or elsewhere, and in its growth produces an enzyme, the secondary infective agent. The enzyme, acting locally upon coagulated albumin, yields soluble albumoses; or being absorbed and conveyed to other organs by the blood, effects a similar change there. The less soluble organic acid mentioned above Martin supposes to be one of the products of the decomposition within the spleen and elsewhere of the albumoses.

Kanthack and Stephens,⁵⁷ who employed ascitic fluid as the albuminous basis of their nutrient substance, observed bacilli in the spleen in a large proportion of fatal cases of diphtheria which they investigated. In the same cases bacilli were also found in the lungs, which were generally the seat of broncho-pneumonia. From the presence of the bacilli in these organs they conclude that the dissemination of the micro-organism by the blood or otherwise is not an unusual event, and object to *Martin's* hypothesis, on the ground that it is consequently superfluous.]

Process for the Detection of the Diphtheria-bacillus.—1. The patient is made to open the mouth as widely as possible, while with a good light, either natural or artificial, a piece of the membrane is removed between the points of very fine forceps. The forceps should previously have been dipped in a boiling solution (1 per cent.) of soda. The shred of membrane thus removed is transferred on a platinum probe to the bottom of the test-tube—both of these being sterilised—and the test-tube is closed with cotton-wool. Pincers and probe are then disinfected by boiling in 1 per cent. soda solution.⁵⁸

2. With similar precautions the specimen is spread upon a slide or cover-glass (p. 46), and stained with Löffler's methylene-blue (p. 47) or with dilute carbol-fuchsin solution.

The specimen, of whatever character it may be, will be found on inspection to consist of fibrinous material, leucocytes and micro-organisms, bacilli and cocci.

Should there appear amongst the latter, bacilli which are slightly curved, about the length of tubercle-bacilli, but somewhat broader,

often a little enlarged at the end, and unequally stained throughout, the extremity being more deeply coloured than the middle part, it is probable that the case is one of diphtheria, pure and simple, and the conclusion is confirmed or otherwise by the clinical symptoms.

The bacilli are often seen to lie parallel to each other in groups, or arranged in angular figures (fig. 50). The appearance presented is highly characteristic, and after some experience it is possible to identify it in cases of uncomplicated diphtheria. The author has repeatedly diagnosed diphtheria in this way, as also have *Heubner* and *Hoppe-Seyler*,⁵⁹ further bacteriological investigations and the clinical developments afterwards proving that his judgment had been correct.

Should the microscopic inspection show only cocci and no bacilli, it is most improbable that the case is diphtheria. If cocci and the bacilli are present together, the conclusion is that the infection is a mixed one.

3. The shred of membrane which was placed in a test-tube, as directed above, is removed on sterilised forceps, washed in 2 per cent. boracic solution (*d'Espine* and *de Marignac*⁶⁰), and implanted on Löffler's blood-serum, solidified obliquely in a test-tube. This is done by stroking the surface of the serum with the membrane.

Löffler's blood-serum has the following constitution: 3 parts sheep's-serum, one part neutralised veal bouillon, 1 per cent. peptone, 1 per cent. grape-sugar, and 0.5 per cent. common salt. *Welch*⁶¹ and many other observers testify to the efficacy of this material in the cultivation of the diphtheria-bacillus. It is a better medium than glycerine-agar, which is presently to be noticed.

[*Kanthack* and *Stephens*⁶² have utilised the serous fluids most easily obtainable in hospital wards as a substitute for Löffler's serum in the preparation of nutrient material for the cultivation of the bacillus of diphtheria. The nutrient substance is prepared as follows:—To 100 cc. of the serous exudation (ascitic, pleuritic, or hydrocele fluid) is added 2 cc. of a 10 per cent. solution of caustic potash, to convert the serum albumin into alkali albumin, and thus prevent precipitation on boiling. To this is added 1.5 per cent. of agar-agar, previously soaked in acidulated water, and the mixture is boiled till the agar-agar is well dissolved, when it is filtered, and 4–5 per cent. of glycerine is added to the filtrate (0.5 per cent. of grape-sugar may also be added, but good results are obtained without this), which is then sterilised in test-tubes.

Its authors claim for this medium considerable advantages beyond the ease with which the albuminous material can be obtained.]

*Toos*⁶³ recommends, as particularly suitable, a nutrient medium containing 20 parts of agar-agar and 20 parts of sodium-albuminate per 1000 parts of peptonised bouillon.

The test-tube containing the infected serum is now placed in a thermostat, maintained at 36.5°–37° C. After the lapse of twelve to fourteen hours there may be seen scattered about the surface of the serum a number of very small transparent points, which, after twenty

to twenty-four hours, grow to the size of a pin's head, and project above the surface, developing at the same time a distinct white hue.

From the cultivation thus made, a cover-glass preparation is obtained in the usual way. If, when examined, this is found to contain only bacilli of the character given above, no doubt remains that the infection was diphtheritic. The author has always adopted this plan when it was merely a question as to the presence of the diphtheria-bacilli.

4. Instead of obliquely solidified Löffler's serum, agar,⁶⁴ to which 6 per cent. of glycerine has been added, may be used, and upon this in a Petri's saucer the membrane is applied after it has been washed with sterilised water⁶⁵ or bouillon.

The nutrient medium adopted as an alternative by Schloffer⁶⁶ consists of 2 per cent. peptone-agar-broth (2 parts), and urine sterilised and heated for half-an-

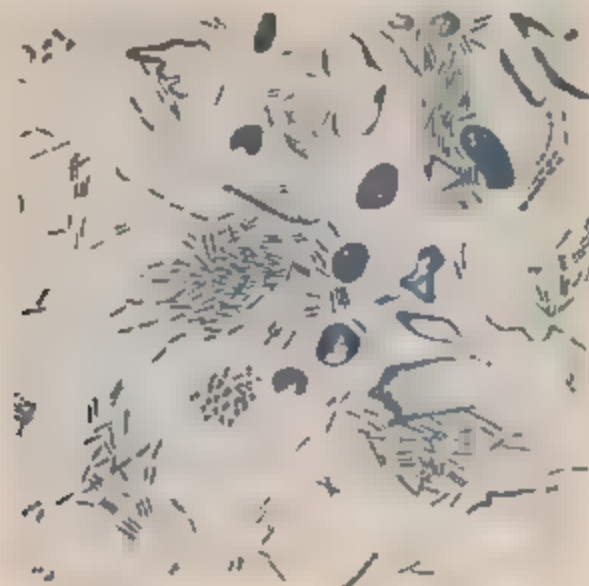


FIG. 50.—Coating of the Tonsils in Diphtheria.

hour at 80° C. This observer maintains that there is not, with any medium, a characteristic development of the broods by which the diphtheria-bacillus may be known.

To inoculate the agar, the shred of membrane is taken in sterilised, still warm forceps of platinum, and the saucer is held over it inverted, that is, with the surface of the agar downwards, while the latter is rubbed over with the membrane between the points of the forceps without breaking the surface, and then the saucer is placed in a thermostat. It is well to conduct the observation at once with two prepared saucers. After eighteen to twenty hours, inspection with a lens of low power (eighty diameters) will show the broods as greyish-yellow, round or oval patches, with ill-defined margin, and liquefied at the edge into an irregular furrow of finely granular fluid. It must, however, be conceded that a cultivation of this kind is not entirely characteristic of

the diphtheria-bacillus. A cover-glass preparation must be made, and if on this are seen only the bacilli (fig. 51) the diagnosis is assured.

When cultivated in alkaline potato the micro-organism forms a greyish-yellow layer, which again is not distinctive.

If, in addition to the appearances just detailed, there are also to be seen on the Löffler blood-serum or agar plate yellowish or whitish patches, which are found by inspection of the dry cover-glass preparation to consist of cocci, the condition is shown to be a combination of diphtheria with infection by Streptococci and Staphylococci. If, again, only the latter are to be seen—as is the case, for instance, in the scarlatina throat—the infection is due to cocci, Streptococci, or Staphylococci, as the case may be.



FIG. 51.—Diphtheria-bacillus (pure culture).

The bacillus of diphtheria has a remarkable vitality. It can be obtained by cultivation from membrane, dried, and eight weeks old.

5. The bacillus is pathogenic in lower animals. In a doubtful instance, and especially to distinguish the bacillus from Hoffmann's pseudo-diphtheria-bacillus, which so closely resembles it, inoculations may be made. Pigeons, guinea-pigs, and rabbits succumb to a small quantity of a pure cultivation, there being first local œdema with fibrinous and haemorrhagic exudation.⁶⁷

6. The diphtheria-bacillus has the property of making neutral bouillon to become alkaline. To display this property a few cc. of nutrient bouillon are taken, a drop or two of litmus solution added, and a pure cultivation of the bacillus implanted on it. After a few days in the incubator the reddish-violet colour gives place to red, and later the cultivation fluid becomes again acid.

The bouillon to be employed is that recommended by *Löffler*⁶⁹ as most suitable for the diphtheria-bacillus. It consists of 2 parts blood-serum and 1 part meat-broth, to which 1 per cent. peptone, 0.5 per cent. common salt, and 1 per cent. grape-sugar have been added.

Baginsky,⁶⁹ and with him the greater number of authorities, holds that the clinical manifestations of diphtheria are produced by two entirely different kinds of micro-organism, and that the clinical type varies accordingly in severity; the more formidable infection being due to Löffler's bacillus, while a milder attack is determined by the admission of Streptococci and Staphylococci within the tissues. The ætiological significance of the bacillus is beyond all question, and it is important to remember that apart from its detection an absolute diagnosis is impossible.

*Peters*⁷⁰ discovered gregarina-like bodies (*Coccidium oriforme*; see chapter on the *Fæces*) in diphtheritic membranes stained with alum-carmine and picric acid. This fact is noteworthy, but further observation must show in what relation these forms stand to the disease in man.

2. Pharyngomycosis leptothricia.—A special interest has recently come to be attached to the nature of the plugs which block the tonsillar crypts. They are found in almost every healthy person, and consist of epithelial cells and of long segmented fungi, which stain bluish-red with the iodo-potassic-iodide solution. In certain conditions these micro-organisms extend outside the follicles, and cover the surface of the tonsils with patches of varying size. They then give rise to subjective symptoms, and their appearance may be mistaken for a commencing attack of croupous or diphtheritic tonsillitis. They may be readily recognised under the microscope by their reaction with the iodine solution, as mentioned above, and the course of the disease will afford a further indication of their nature (*Th. Hering*).⁷¹ *O. Chiari*⁷² is of opinion that this affection is not one *sui generis*, but should merely be regarded as a modification of *Angina follicularis*, in which such products are always found.

One or two minutes in the iodine solution suffice to develop the bluish-red colour in a specimen containing leptothrix. This colour disappears in from twenty-four to seventy-two hours.

Dr. O. Chiari has called the attention of the author to the fact that yellowish plugs which do not contain leptothrix are often to be found in the crypts. In a very hard concretion from the tonsils, found on chemical examination to consist of carbonates and silicates, the author met with splendid specimens of leptothrix.

CHAPTER III

THE NASAL SECRETION

I. NAKED-EYE AND MICROSCOPICAL CHARACTERS—

CHEMICAL CONSTITUTION.—Considering the great quantity of glandular tissue with which the nasal passages is furnished, the secretion in health is remarkably scanty.

Normal nasal mucus exhibits microscopically squamous and ciliated epithelium in abundance, isolated leucocytes, and an enormous profusion of fungi.

*E. Weibel*¹ has described a curved bacillus obtained from the nasal secretion of healthy persons, which, when cultivated in nutrient gelatine

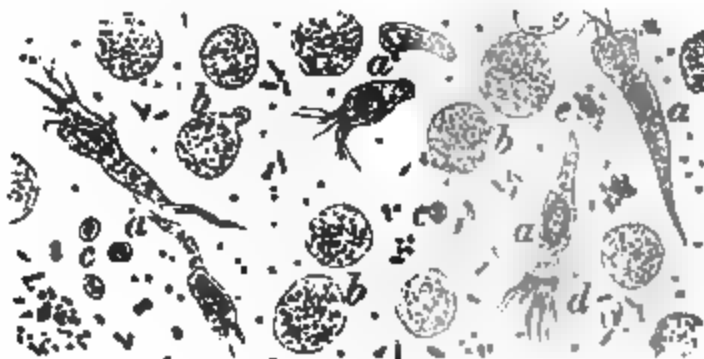


FIG. 52.—Nasal Mucus (eye-piece III., objective 8*A*, *Reichert*).
a. Ciliated epithelium; b. Leucocytes; c. Encysted cocci; d. Bacilli; e. Micrococci.

and agar-agar, develops a spirillum-like body wound into several coils. It is probable that further researches will show the presence of many other forms. A great variety has been enumerated by *Reimann*.²

The normal nasal secretion is a thick fluid, faintly odorous, and of an alkaline reaction. It abounds in mucin, but otherwise nothing definite is known about its chemical constitution.

II. THE SECRETION IN AFFECTIONS OF THE NASAL CAVITIES.—At the outset of an attack of acute nasal catarrh, the mucous membrane is generally dry and much injected, and the secretion lessened

in quantity. Later on, however, there is a copious discharge of a thin alkaline fluid, and this, when examined under the microscope, is seen to consist of a great number of epithelial cells and fungi.

Where suppuration is in progress within the nose, the secretion partakes of the character of pus, and is seen microscopically to consist almost entirely of pus cells. Occasionally, as in cases of wounds perforating the cranium and in brain tumours, cerebro-spinal fluid may be discharged through the nose. *Nothnagel*⁸ has reported a very interesting case of this kind. Under such circumstances, chemical analysis showing the absence of albumin and the presence of sugar, or, at least, of a reducing substance, will determine the diagnosis. The importance of the evidence so obtained in connection with cerebral tumours is evident.

It is very necessary, in all cases of ulceration of the mucous membrane of the nose, to look for certain of the pathogenic fungi already known to us.

Thus, if it be a question whether a particular ulcer is tubercular or not, a little of the discharge may be removed with the help of the nasal speculum on a carefully sterilised platinum spatula, and examined for tubercle-bacillus in the manner indicated at p. 46.

According to *Sticker*⁹ the examination of the nasal secretion for lepra-bacilli is of great importance in suspected cases of leprosy, which complaint he regards as primarily a nasal disease.

Again, the discovery of the characteristic bacillus of glanders in such a discharge will obviously determine the diagnosis of that disease. This bacillus may be sought for in the same manner as in the examination of blood (p. 52). If this be not enough, fungi may be cultivated on *Koch's* plan (see Chapter X.) ; or, finally, in case of doubt, they may be propagated in one of the lower animals, when a definite conclusion will be arrived at.

*E. Frankel*⁵ and *Hajek*⁶ observed various forms of fungi constantly present in the discharge of the chronic ulcerative processes known as ozæna. *Löwenberg*,⁷ on the other hand, found that one large species of diplococcus was almost the only form present in such discharge, and he regards it as characteristic of ozæna.

*Tost*⁸ and *Lorenberg*⁹ have shown that bodies resembling the pneumonia-coccus occur in the nasal secretion (fig. 52, c). *Abel*¹⁰ found a bacillus in sixteen cases of simple ozæna. *H. v. Schrötter* and *Winkler*¹¹ have isolated the *Staphylococcus cereus flavus* and another similar micro-organism, which they designate *albus*, from the nasal discharge in coryza. The discovery is obviously of no importance unless it can be shown that these forms do not occur in health.

Rhinitis fibrinosa may here be specially mentioned. While some

authors regard this as an affection distinct from diphtheria,¹² the observations of others¹³ show that it may be induced by the diphtheria-bacillus. It is evident that under the term Rhinitis fibrinosa various distinct diseases are included. The striking observations of *Mya, Gerben, Podack, v. Starck*, and others,¹⁴ bear upon this point.

In a few cases, thrush-fungus and vegetations have been found in the nose. Mould-fungi in this situation are another rare manifestation (*Schubert*).¹⁵ Ascarides and other entozoa are very seldom seen there. *Proskauer*¹⁶ believes that he has found the embryo of *Oxyuris*. The statement is very doubtful. Dipteral larvæ are of most common occurrence (*B. Fränkel*).¹⁷

The *Charcot-Leyden* crystals found in blood and sputum have also been met with in the nasal secretion of an asthmatic patient. *Leyden*¹⁸ observed them associated with eosinophil cells in the nasal mucus in a case of acute coryza, and *Sticker*¹⁹ in blood discharged from the nose in leukæmia, after it had stood for some days. *Levy*²⁰ has described them in connection with nasal tumours (polypi).

Concretions (rhinoliths) occasionally form in the nasal cavities (*O. Chiari, Seifert*).²¹

CHAPTER IV

THE SPUTUM

UNDER the term *expectoration* or *sputum*¹ are comprised all those substances which are removed from the air-passages by the mechanical effort of coughing or hawking.

1. NAKED-EYE CHARACTERS OF THE SPUTUM.—The naked-eye appearance of the sputum will often afford valuable information ; and that this may be as accurate and exhaustive as possible, it is well to collect the expectoration in glass vessels, after *Nothnagel's* plan, and then to examine it as to its quantity, specific gravity, reaction, colour, smell, and tendency to stratification.

The quantity of expectoration discharged in twenty-four hours varies within broad limits. Sometimes it does not exceed a few cc. ; in certain conditions, on the other hand, as, e.g., where an empyema is discharging into the lung, as much as 800 to 1000 cc. may be expectorated in twenty-four hours.

*H. Kossel*² determines the specific gravity of the sputum in the following manner :—The sputum is placed in a flask stoppered to prevent evaporation, and gradually heated to 60° C. It is thus reduced to a thinly-fluid state, and is placed in the pycnometer. The specific gravity is seen to vary within very broad limits,—for mucous sputum, 1.0043–1.0080 ; for purulent, 1.0155–1.0260 ; and for serous, 1.0375. The question of density is of only slight interest in clinical cases.

The reaction of the sputum is always alkaline.

In some diseases, as in abscess and gangrene of the lung, there is marked *stratification* of its parts (see p. 147).

The colour of the sputum depends partly upon its microscopical and partly upon its chemical character. When it consists entirely or chiefly of mucin and a few cells, it is whitish. Green sputa are usually purulent, but the presence of biliverdin or of pigment-forming bacteria may also impart this colour.

The odour of the sputum is, for the most part, not characteristic ; but in putrid bronchitis and gangrene of the lung it has a particularly pungent and unpleasant smell.

For many purposes it is convenient to collect the sputum in a cylindrical glass vessel containing water. In this way, for instance, the nummular arrangement is made apparent. In other cases, again, certain constituents, as, *e.g.*, spirals, fibrinous coagula, and shreds of tissue, may be made more evident on a dark surface, such as that of a polished black plate. A very suitable plate has been devised by *Kroenig*.³

*Ad. Schmidt*⁴ has employed Ehrlich's triacid mixture (see p. 36) with success in the investigation of the sputum.

A passing reference must be made to the staining methods of *Schmidt* and *Lilienfeld*.⁵

[The expedient of hardening the sputum and examining it in sections was first adopted by *Ad. Schmidt*, and lately revived by *Gabritschewsky*.⁶ As fixing and hardening fluids the latter employs alcohol, Müller's fluid, Flemming's solution, picric acid, and sublimate in concentrated forms. The hardened sputum is embedded in celloidin for cutting, and the sections stained. This method is valuable for the detection of some of the constituents of the sputum, which may be destroyed by pressure under a cover-glass.]

Although much knowledge of a disease may be derived from a naked-eye inspection of the sputa, it will never enable us to dispense with the aid of the microscope, by means of which alone we can diagnose certain affections—some forms of tuberculosis, for instance—with the utmost certainty.

II. MICROSCOPICAL EXAMINATION OF THE SPUTUM.

1. White Blood-Corpuscles.—These bodies are always found in large numbers in the sputum, commonly embedded in a viscid stringy substance. Many of them are of large size and granular, and enclose within them drops of fat and particles of pigment, such as carbon dust and masses of hæmatoidin (see fig. 53, *e*). Eosinophil granules are found in the leucocytes of the sputum in certain bronchial affections. In cases where an abscess has discharged into the lung, and in purulent bronchitis, such as is met with in connection with emphysema, the sputum may consist entirely of leucocytes.

[Eosinophil granules have been observed in the sputum of phthisis (*Janowsky*),⁷ asthma, (*Gollasch*),⁸ and bronchitis (*Kanthack*).⁹]

Attention has recently been drawn by *F. A. Hoffmann*¹⁰ to the presence in sputum of certain cells, which he terms "Schläuche" (hose-pipe cells). These, he believes, occur mainly in acute processes in the sputum.

2. Red Blood-Corpuscles.—These are also to be found in almost all sputa, and their presence in small numbers is without significance. In persons who smoke a great deal, or who spend much of their time in an atmosphere of tobacco-fumes, the sputa are apt to be streaked with blood in the morning. This blood, however, proceeds in most cases,

not from the lung-tissue proper, but from the bronchial mucous membrane, and is due to catarrh.

When red blood-cells are present in very considerable quantity, the sputa will be coloured by them. The individual cells are usually intact, and in this they depart from the condition in which they are found in urine and faeces. In some cases, however, as in pneumonia, they are altered and occur as pale discoidal bodies. Where blood has accumulated for some time in the bronchial tubes, the red corpuscles may disappear entirely, and the pigment remain in the sputum either as irregular particles or as red crystals of haematoxin (see fig. 53, e). The sputum of pneumonia derives its colour from blood pigment dissolved in it.

Finally, it is to be noted that in haemorrhage from the lungs the

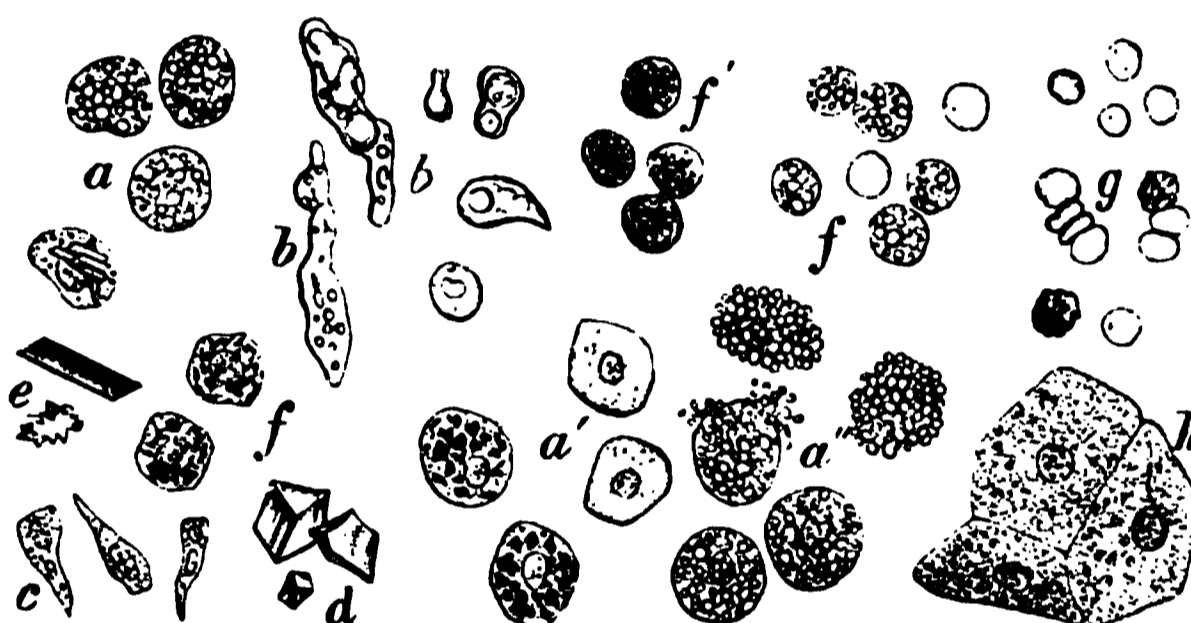


FIG. 53.—Epithelium, Leucocytes, and crystals of the Sputum (eye-piece III., objective 8A, *Reichert*).

a, a', a''. Alveolar epithelium. *b.* Myelin forms. *c.* Ciliated epithelium. *d.* Crystals of calcium carbonate. *e.* Haematoxin crystals and masses. *f, f', f''.* White blood-corpuscles. *g.* Red blood-corpuscles. *h.* Squamous epithelium.

sputum consists entirely of red corpuscles; in congestion the blood is intimately mixed with mucus.

3. Epithelium.—The sputum abounds in epithelial cells.¹¹ Squamous cells (fig. 53, *h*) come either from the mouth or from the surface of the true vocal chords. Ciliated epithelium is less often seen, and occurs chiefly in severe bronchial inflammation, when, too, it is probably derived rather from admixture with nasal mucus¹² than from the surface of the trachea, which, as is well known, is lined with ciliated epithelium. The cells as found in the sputum are usually deprived of their cilia (fig. 53, *c*), unless in quite recent expectoration, when cilia in active motion may still be seen on them. The mere presence of such cells in the sputum is of little diagnostic import; but where they occur in great numbers they may be taken as indicating the commencement of acute catarrh, either in the back part of the nasal fossæ or in the trachea and bronchi.

But there is another variety of epithelium whose appearance in the sputum is a fact of great importance.¹³ It is known as "alveolar" epithelium, a name which will serve provisionally, although its derivation from the alveoli of the lungs has lately been called in question (*Bizzozero*).¹⁴ It consists of elliptical cells, each containing one nucleus, which usually requires acetic acid to make it visible. The body of the cell is of finely granular protoplasm, and very often holds irregular pigment particles in its substance. These particles consist usually of blood-colouring matter, iron dust or carbon (fig. 53, *a'*). In the last case they are unaffected by reagents generally. Iron dust may be known by its turning blackish-green with sulphide of ammonium, or blue with yellow prussiate of potash and hydrochloric acid. These cells often contain one or more fatty granules, readily recognisable by their high refractive power, and at times they exhibit extreme fatty degeneration (fig. 53, *a, a''*), when their protoplasm is replaced by oily drops of varying size. Sometimes large bodies like drops of fat, presumably derived from the rupture of these cells, are to be seen in the sputum (fig. 53, *b*). They were first observed by *Virchow*,¹⁵ who named them *myelin droplets*, from the resemblance they bear to similar forms produced in the destruction of nerve tissue. According to *Panizza*,¹⁶ however, this myelin (which is only the outward form of a considerable number of different substances) is probably mucin, while *Zoja*¹⁷ suggests it is lecithin or a protagon; but no diagnostic importance is attached by them either to myelin or to myelin-holding cells.

*Buhl*¹⁸ thought that the appearance of alveolar epithelium in the sputum was characteristic of the disease which he has named desquamative pneumonia. It is certainly a fact that such cells are to be found in great profusion only in quite fresh specimens of caseous infiltration of the lung, whether due to bacilli or not; but then they occur also in pneumonia, in chronic bronchitis, and in chronic pulmonary tuberculosis;¹⁹ sometimes, too, in very large numbers. It follows from their manifestation in processes differing so entirely that their diagnostic significance is on the whole slight.

[*Troup*²⁰ believes that the presence of alveolar epithelium in the sputum belongs especially to obstinate catarrhs of the apex, when it will be found associated with columnar and ciliated cells; he observes that since such catarrhs tend for the most part to run into phthisis, we have in this an early and valuable sign of impending danger.] There is a special form of alveolar epithelium, consisting of large flat cells containing a golden yellow and brown pigment (failing-heart cells, *Wagner*), whose presence in the sputum, according to *F. A. Hoffmann*,²¹ only occurs in valvular heart-disease and adherent pericardium. They are absent from the sputum in phthisis and pneumonia, and they may

therefore be taken, in doubtful cases, to point to brown induration of the lung. Whilst agreeing in general with Hoffmann's view, the author would insist that in the processes just mentioned the sputum exhibits cells containing black pigment, which chemical examination has shown to be a derivative of blood-colouring matter. Such cells were noticed long ago by J. Sommerbrodt,²² who described them as brown alveolar epithelium. He is in accord with Hoffmann as to their meaning. Lenhartz²³ has observed them most frequently in connection with mitral stenosis, and regards them as metamorphosed blood-corpuscle-holding round cells (lymph corpuscles). Kronig²⁴ endorses Hoffmann's view; and v. Noorden²⁵ believes the cells to be pigment-bearing eosinophil leucocytes.

To examine the sputum for epithelium, a small quantity should be

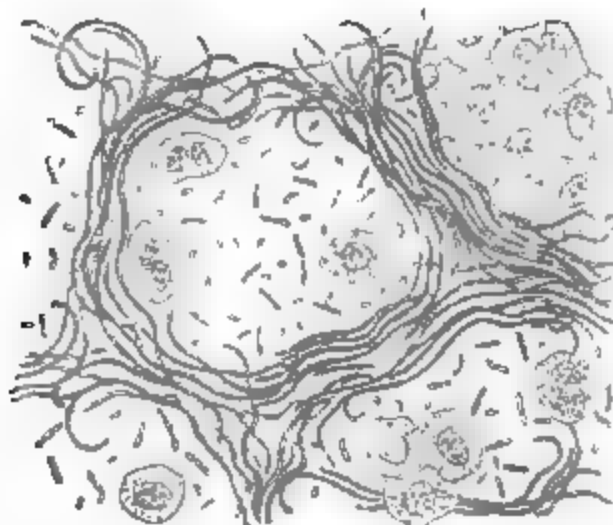


FIG. 54.—Elastic Fibres in the Sputum (eye-piece III., objective 8A, Reichert).

treated with acetic acid, to bring the nuclei and nucleoli into view; or a specimen may be stained with a watery solution of methylene blue.

4. Elastic Fibres.—These fibres occur in the sputum singly or in bundles, and they have commonly an alveolar arrangement (fig. 54). They are of varying length and breadth, dark-coloured, slightly curved, and generally exhibit a double contour.* Their diagnostic value is great, as a sign of serious mischief, pointing to destruction of lung-tissue. They occur, consequently, in tuberculosis, bronchiectasis, pulmonary abscess, and occasionally in pneumonia, while the other signs of abscess are wanting. The author has repeatedly found elastic fibres in cases of pneumonia which otherwise ran a favourable course, and he supposes that in such cases there was destruction of the pulmonary parenchyma only over a very limited area. It is a notable fact that

* The figure is taken from a case of pulmonary abscess which was for several months under treatment in Professor Nothnagel's clinic.

these fibres are rarely to be met with in the expectoration of pulmonary gangrene, and the reason probably is that they are destroyed *in situ* by the ferments formed in that process. The fact was first noticed by Traube.

Elastic tissue may be introduced with the food, and so find its way into the sputum. A necessary precaution, therefore, is to direct the patient to wash the mouth carefully after food, and to separate the sputa discharged at meal-times from those to be examined for lung-tissue. Even when every care has been taken, however, one is exposed to fallacy arising in this way, as fibres derived from the food may lie in the mouth for days before they are expectorated. *It is only when the bundles of elastic fibres display the alveolar arrangement that we can be certain of their origin in the pulmonary alveoli, and it is then only that they possess any sure diagnostic significance.*

For their detection, when they occur in considerable quantity, it will suffice to add some potash solution to a little of the sputum placed on a

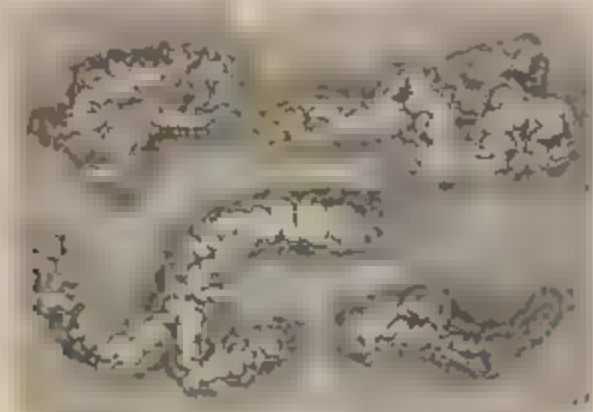


FIG. 55.—Spirals from the Sputum, in a case of bronchial Asthma (actual size)

slide. A still better plan is that proposed by Fenwick, who boils the sputum in a solution (8-10 per cent.) of caustic potash, allows the mixture to stand for twenty-four hours in a conical glass, and examines the sediment for elastic fibres under the microscope.

[Prolonged boiling in caustic potash will cause the elastic fibres first to swell up and then to dissolve. Consequently the boiling must not be too long, and Dr Troup recommends that when the sputum is placed on a slide the cover-glass should remain undisturbed.]

5. Spirals.—Certain spiral bodies were first observed by Leyden²⁶ in the sputum of patients suffering from asthmatic paroxysms.

They were afterwards described by Curschmann²⁷ as pathognomonic of disease of the finest bronchial tubes; and since then they have been seen by O. Vierordt, v. Jähn, Pet, and Sanger²⁸ in the sputum of pneumonia. Recent personal observations have satisfied the author that

these bodies are also found in the sputum in cases of acute pulmonary tuberculosis. More recently *Lewy*²⁹ and *Troup*³⁰ have written on their occurrence in asthmatic paroxysms.

*Korotz*³¹ has seen them in pulmonary oedema, and *Czermak*³² observed quite similar bodies in vascular keratitis. He believes that they are produced by a twisting upon their axis of the ordinary mucous threads, and he succeeded in forming well-shaped spirals experimentally by twisting them in this way. *V. Gerlach*³³ arrived at the same conclusion. *R. J. Jaksch*³⁴ has described similar bodies as seen in the urine and faeces.

The spirals are usually discernible with the naked eye when the sputum is carefully examined. They appear then as thick white bodies of a twisted and tubular form, and distinguished by their clear tint and great tenacity from all other sputum constituents (fig. 55).

Under the microscope they display a remarkable variety of shape. In



FIG. 56.—Spirals from Sputum (eye-piece III, objective 4, *Steinert*).

the usual arrangement there is a central thread disposed in a more or less zigzag manner, and around it a thick meshwork of very delicate fibres commonly looped round in spirals, but occasionally retiform. The spirals are often overlaid with epithelium, and sometimes also covered with *Charcot-Leyden* crystals. They vary greatly both in length and breadth. When present, they indicate so it would seem—a desquamative catarrh of the bronchi (*Curschmann*) and alveoli (*Lewy*). Thus they are found in pneumonia as well as in capillary bronchitis. In connection with asthma they afford a most valuable diagnostic sign, showing that the disease is of a bronchial character.

As to the connection between these spirals and the *Charcot-Leyden* crystals (see p. 34 and Chaps. VI and IX.), it is to be noted that in the earlier attacks of bronchial asthma, or at the beginning of a fresh attack, spirals are to be found in the sputa, but no crystals. If, however, a

specimen of such a sputum is placed quite fresh on a slide covered to prevent evaporation, and allowed to stand for 24-48 hours, crystals will be seen in it. Later in the asthmatic paroxysm, numbers of similar crystals occur in the recent sputa, lying amongst the spirals, and not elsewhere. From this it would appear that the *Charcot-Leyden* crystals are in part derived directly from the spirals. As to the chemical constitution of the latter, the author has ascertained that they are formed of a substance closely resembling mucin. If one of them be removed from the sputum and touched with a dilute alkali, it dissolves; and when acetic acid is added a precipitate forms. If the alkaline solution be treated with cupric sulphate and boiled, the resulting peroxide of copper is not reduced; but with the previous addition of a mineral acid and boiling, reduction immediately takes place. All these reactions are common to mucin.

Observations which the author had recently an opportunity of making



FIG. 57.—Fibrinous Coagula from a case of Pneumonia, with Paroxysms of severe Dyspnoea (I).

in a case of bronchial asthma went to confirm the statements made above, and showed further that the portion of the spirals constituting the central thread is chemically distinct from its envelope of mucin. This central thread approached rather to the character of fibrin, but its precise composition was not ascertained with certainty.

Asthmatic sputa contain large numbers of leucocytes containing eosinophil granulations (*Fr. Müller, Gollasch, Schmidt*).³⁵ The fact, however, is of little aid in diagnosis, since *Gollasch* and *Leyden*³⁶ have observed similar bodies in the expectoration of acute and chronic bronchitis.

6. Fibrinous Coagula.—Fibrinous casts are found in the sputa of plastic bronchitis and pneumonia. They are whitish in colour, of varying consistence, and branched to the form of the terminal bronchial tubes in which they are deposited. In pneumonia they are usually few in number and of no great length (fig. 57), and when more numerous they mark a severe clinical type of the disease. The abundant formation

of such coagula in pneumonia is attended with much coughing and great dyspnoea.

They are found in greatest perfection in the chronic plastic bronchitis of adults. They may be said to be pathognomonic of this affection, which is otherwise so often difficult to diagnose. The author has seen them several centimetres long (fig. 58), and minutely branched. Microscopically these terminal filaments form a fine network, in which epithelial cells and blood-corpuscles are enclosed. These coagula being formed of fibrin, resist the action of acetic acid. Their chemical investigation may be conducted in the manner described under *Fibrinuria* (Chapter VII.).

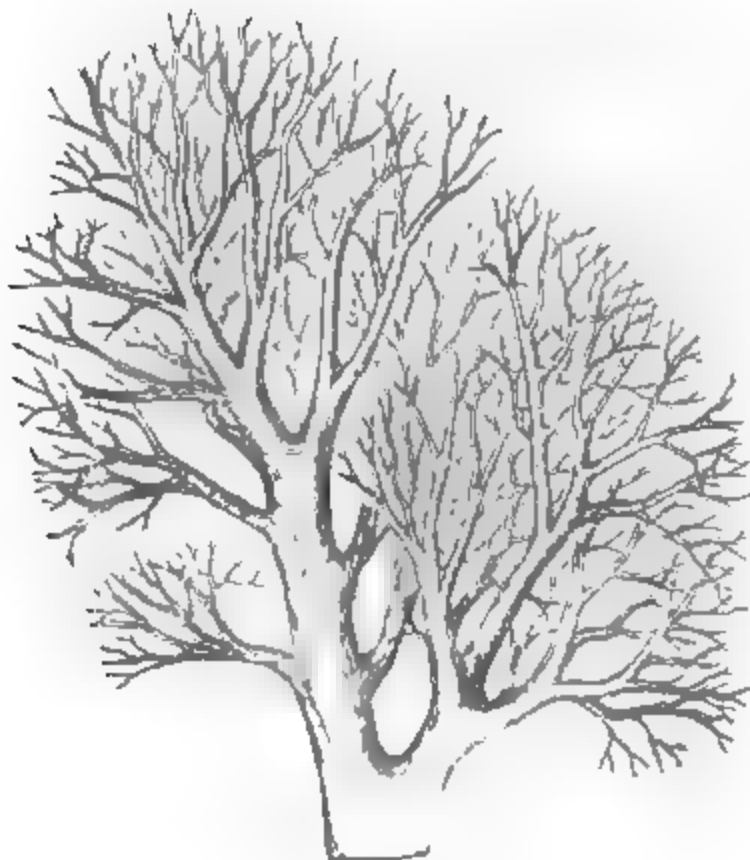


FIG. 58.—Fibrinous Coagula from a case of Plastic Bronchitis (J).

The author, having regard to the resemblance clinically between chronic plastic bronchitis and bronchiolitis exudativa, has examined the sputum in such a case for eosinophil cells, but without success.

7. Connective Tissue.—Shreds of connective tissue are occasionally, though rarely, found in the sputum. It sometimes happens in pulmonary abscess and gangrene that scraps of tissue are coughed up, which may be recognised by their arrangement as belonging to the alveoli of the lungs. Fragments of cartilage, due to ulceration of the larynx, may also be discharged with the expectoration. Their origin can usually be determined by the microscope. In sarcoma of the lung particles of tissue

are apt to be discharged, and these, by their characteristic appearance, enable a definite diagnosis to be made (*Huber*).³⁷

8. Corpora Amylacea.—*Friedreich*³⁸ has described these starch-like formations as occurring in the sputum. He refers their origin to haemorrhagic processes in the lungs. They are partly round, partly angular in shape, and contain within them a core of pigmentary substance, which also is usually angular. In a solution of iodine and iodide of potassium, they sometimes give the amyloid reaction, but not always. Often their form shows stratification. In a sputum sent to him for examination by his colleague, *Dr. Neusser*, the author found such bodies, and he has several times met with others like them, but without the dark central substance in the sputum of pulmonary gangrene. These did not give the amyloid reaction, but they showed evident stratification. It must still be regarded as an open question whether these bodies are of an amyloid nature or not.³⁹

9. Parasites.

1. Fungi.—The investigation of the sputum for fungi has in recent times engaged much of the attention of scientists and physicians. If we adhere to the usual classification of parasitic fungi into moulds, yeasts, and fission-fungi, it will be found that the chief interest attaches to micro-organisms of the third class. Of the fission-fungi some indeed are innocuous; but there are others of a distinctly pathogenic character, and their recognition in the sputum is a point of prime importance in diagnosis. It will serve a useful purpose, therefore, if we divide the micro-organisms in question into two chief classes, the *non-pathogenic* and the *pathogenic*, bearing in mind at the same time that parasites which are ordinarily harmless may at times be the cause of the most formidable diseases, whilst, on the other hand, certain others that are under appropriate circumstances the most dangerous pests of humanity, as, for instance, the *Diplococcus pneumoniae* and the diphtheria-bacillus, occur also in the sputum of quite healthy persons.

(a.) Non-Pathogenic Fungi.

i. Moulds.—There is little to be said of these parasites as a constituent of the sputum. The thrush-fungus rarely occurs (see p. 102), and its presence is usually to be referred to admixture with the saliva. In all cases where it is found, the mouth and throat should be carefully examined for aphthæ. At the same time it must not be forgotten that in thrush aphthous patches may extend to the bronchi, especially where the subjects are children.

Further, in certain pulmonary diseases mould-fungi have been found in the sputum. The accompanying figures (fig. 59) show such bodies as seen in the fresh sputa of a patient suffering from traumatic abscess of the lung. *Virchow*⁴⁰ has published observations bearing upon this

subject, and *Lichtheim*⁴¹ found *Aspergillus fumigatus* in human sputa. This latter organism has been shown by *Schütz*⁴² to be connected with disease in the lower animals. *Coppen Jones*⁴³ also has described a mould-fungus which occurs in the sputum of phthisis.

[*Arkle* and *Hinds*⁴⁴ have recorded a case of extreme emphysema, occurring in a young man otherwise healthy, and whose conditions of life excluded the ordinary causes of pulmonary disease. The lungs were found at the autopsy to be permeated with a mycelium believed to be *Aspergillus*.]

Most writers have concurred in the opinion that mould-fungi are only

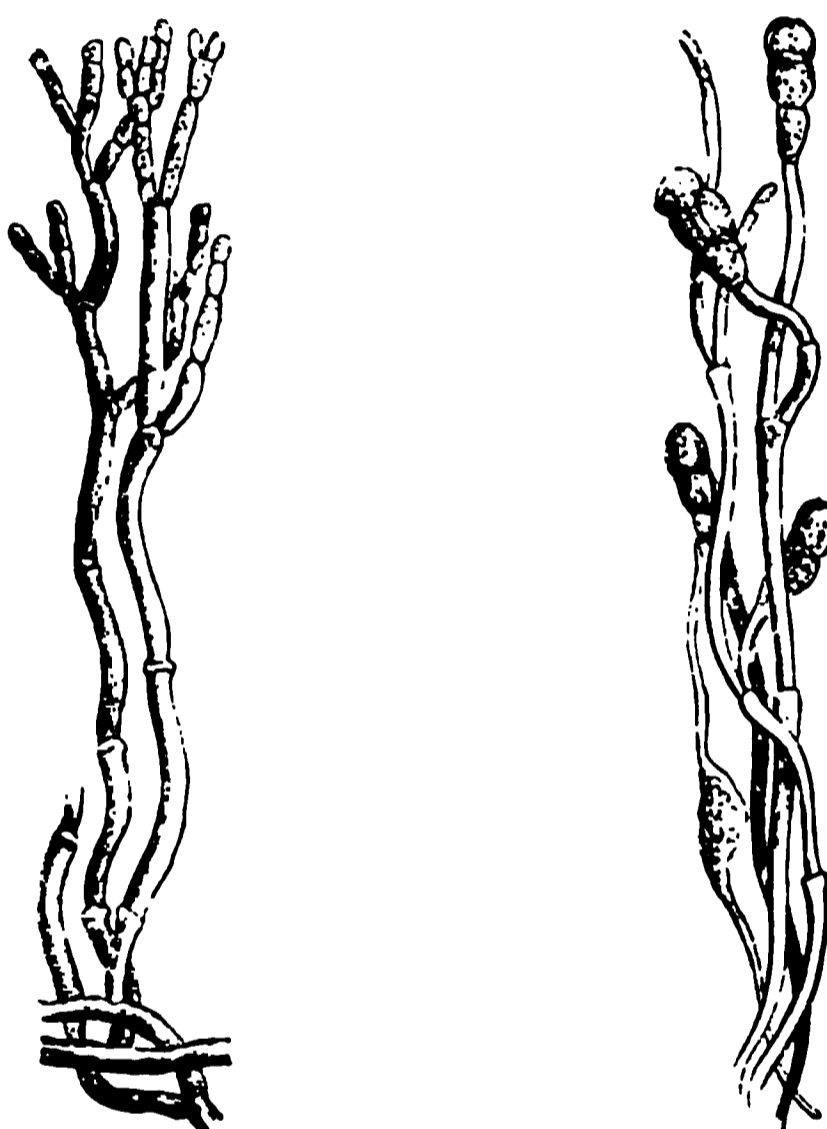


FIG. 59.—Moulds from the Sputum in Pulmonary Abscess (eye-piece I., objective 8A, *Reichert*).

incidentally present in the sputum; but quite recently the labours of *Schütz*, based upon the result of *Lichtheim's* experiments with animals, have gone far to show that the multiplication of mould-fungi alone may cause destructive processes in the lungs. The observations of *Paltauf* and *Lindt*⁴⁵ tend to the same conclusion. To study the nature of these fungi they should first be found by microscopical examination, and then cultivated on bread and nutrient gelatine. Their relation to disease may also be ascertained by experiments upon animals (see Chapter X.). Whether innocuous or not, it is quite certain that mould-fungi are frequently present in the expectoration.

2. **Yeasts.**—Little is practically known of yeast-fungi as constituents of the sputum. The author has discovered scattered yeast-cells in the pus from a phthisical cavity.

3. **Fission-Fungi.**

1. **Sarcina Pulmonis.**—Sarcinæ have been found in the expectoration of several diseases. They were first described in this connection by *Virchow* and *Friedreich*.⁴⁶ They are usually smaller than *S. ventriculi*.⁴⁷ Their presence is of little pathological interest (*Fischer, Hauser*⁴⁸), but it would seem that they belong especially to extensive ulceration of the lung. *Pansini*⁴⁹ describes a new species under the name *Sarcina variegata*.

2. **Leptothrix.**—*Leyden* and *Jaffé*⁵⁰ have repeatedly found leptothrices in the sputum (see p. 103). They may be recognised in the so-called fungoid plugs of putrid bronchitis by their property of staining in iodo-potassic-iodide solution. These fungoid plugs have been analysed by *Dittrich, Traube*, and others. They contained, besides leptothrices, crystals of hæmatoidin, white and red blood-corpuscles, and in many cases quantities of fat-laden epithelial cells and a mass of fatty detritus.

3. **Bacilli and Micrococci.**—Various forms of bacilli and micrococci occur in every sputum. A number of these, and among them bacilli with terminal spores, are shown in fig. 54.

(b.) **Pathogenic Fungi.**

1. **Tubercle-Bacillus.**—The researches of *Robert Koch*⁵¹ have shown that the sputum from tubercular lungs contains definite organisms which behave in a uniform and remarkable manner towards certain staining fluids; and this author was further of opinion that the organisms in question were the vehicles of the tubercular virus. His conclusions have been most fully borne out by subsequent observations.⁵² Nothing more need be said just now to show the great importance of his discovery. We shall return to the subject later when we come to speak of tubercular sputum.

The tubercle-bacillus cannot be seen in the unstained sputum. When a preparation is made in the manner to be described presently, the bacilli appear either separately or more usually in groups, as small rod-like bodies, slightly curved, very delicate, and of variable length ($1.5 \mu - 3.5 \mu$). They are quite motionless, and spore-formation can sometimes be seen in them. The spores do not stain by the same process as the parent-bacilli, and thus it happens that the tubercle-bacillus will seem to be broken up into several (2-6) clear oval compartments. This appearance has led some observers⁵³ to suppose that they had to do with micrococci, but by appropriate methods and careful examination the clear outline of the bacillus can be made visible in such cases through its entire length (fig. 60).

Detection of Tubercle-Bacillus.—Since the tubercle-bacillus was discovered some years ago, very many methods have been suggested for its recognition, as those of *Koch*, *Ehrlich*, *Gibbes*, *Baumgarten*, *Neelsen*, *Balmer*, *Fräntzel*, *Kühne*, *Fränkel*, and *Gabbett*. They all alike depend upon the remarkable property which the bacillus displays of *staining with aniline dyes in alkaline solution, and (unlike the other micro-organisms, pathogenic and non-pathogenic, which occur in the sputum) of retaining the dye in after-treatment with acid and alcohol.*⁵⁴

The bacillus may be shown by any of the methods enumerated above when the requisite skill has been attained, but the methods of *Koch* and *Ehrlich* are the best for a beginner to employ.

A. *Preparation of the Staining Fluid.*—It is important that the staining fluid should be freshly prepared for use in every case, both because the aniline dyes are apt to undergo chemical changes when kept for a time in solution, and also because fungi may develop in the fluid itself, and so mislead the observer. It may be made thus: In a test-tube, carefully washed (with distilled water and alcohol) and dried, an alcoholic solution of gentian- or methyl-violet is made * by dissolving the solid substance in 4–5 cc. of absolute alcohol, and to such a point of concentration that an object cannot be discerned through the fluid in the test-tube.

In another test-tube, which also must be thoroughly cleaned, 6 cc. of distilled water are measured out; to this are added 10–15 drops of aniline oil. The fluid is well mixed by shaking it up, and then passed through a moist filter. To the clear filtrate are added a few drops of the alcoholic solution of gentian- or methyl-violet, prepared as above, till a slight turbidity appears. This should clear in a few minutes. Should the mixture fail to become perfectly clear, however, that will not interfere with the success of the undertaking. Thus is obtained the aniline-water-gentian- or methyl-violet solution (*Weigert-Ehrlich fluid*). In addition a watery solution of Bismarck-brown or vesuvin is needed. To make this, a small quantity of one of these substances (about as much as could be taken up on the point of a pen-knife) is added to a few cc. of distilled water in a test-tube, and dissolved until the fluid is barely transparent; it is then filtered. The filtrate will be employed in the manner to be indicated presently.

B. *Preparation of the Cover-Glasses.*—The cover-glasses to be used must be scrupulously cleaned by washing first in distilled water and then in strong alcohol, and afterwards dried in an exsiccator, or at any rate in a chamber free from dust. A number of such thoroughly cleansed cover-glasses should be kept ready for use under a glass shade.

* To simplify the method, a concentrated alcoholic staining solution might be kept in readiness.

One is now taken up with forceps, previously sterilised by raising them to a white heat, and with similar forceps a particle of the sputum to be examined is placed on the cover-glass—such portions as appear most purulent being chosen—and spread in a layer as far as possible of uniform thickness. Another cover-glass is placed upon the first, and the two are pressed closely together with the forceps, so that the inclosed sputum is spread very thinly. The cover-glasses, still grasped with the forceps, are then separated, and dried first in the air, and then passed two or three times through a carbon-free flame from a spirit- or gas-lamp, the preparation surface being kept upwards. *V. Rindfleisch*⁵⁵ advises that a camel's-hair brush be used, this being dipped in the sputum, and the cover-glass painted with it. The thin fluid thus obtained is particularly rich in bacilli. The same brush must not, of course, be used a second time. [When a film is made in this way, cover-glasses can be dispensed with. The secretion is then applied directly to a slide, and the preparation stained in a bath. When dried, the preparation is covered with cedar oil, in which also the lenses are immersed.]

C. *To Stain and Mount the Preparation.*—The cover-glasses, prepared in the manner just described, are now placed in a watch-glass holding the aniline-water-gentian-violet staining fluid already mentioned. They should be allowed to float in the fluid preparation-side downwards. When removed after twenty-four hours they will be found to be coloured a dark blue. They must now be placed in dilute nitric acid (1 in 4) until the preparation appears to the naked eye no longer blue, but at the most a bright green.

The preparation should not be completely decolorised when taken from the fluid, because there is danger that the bacilli also may be deprived of the stain if they are submitted to a too prolonged action of the acid. When removed from the latter, they must be rinsed with absolute alcohol, then dried in the air, and mounted in oil of cloves or Canada balsam.

For microscopical examination Hartnack's objective No. VII., or Reichert VIII., serves best with a little practice. The beginner may use an oil-immersion lens and Abbe's condenser with advantage. If tubercle-bacilli are present, they will appear in the specimen as minute rod-like bodies of a blue colour, but if few in number they may easily be overlooked. It will then promote their detection to stain the surrounding substances by immersion in the solution of Bismarck-brown or vesuvin. To do this the cover-glass, prepared and stained in the manner already described, is placed in the brown staining fluid (made as above), and left in it until the whole preparation has assumed an evident brownish-yellow tint. It is then washed with distilled water and mounted for examination. When looked at through the microscope,

the bacilli are seen to be coloured a deep blue, while the other bodies in the field—fungi, mucus, corpuscles, and epithelium—are brown (fig. 60). The length of the individual bacilli as found in the sputum is very variable. The author has repeatedly met with specimens as large as those represented in fig. 60, but smaller forms, as shown in fig. 61, occur still more frequently. The entire process may be accomplished in a quarter of an hour by heating the staining fluid, which also should then be a concentrated solution of the aniline-water and gentian-violet mixture.

The method of staining with carbon-fuchsin (*Ziehl-Neelsen* fluid⁵⁶) is well worthy of notice here. Ten cc. of a concentrated alcoholic solution of fuchsin are added to 90 cc. of a 5 per cent. solution of carbolic acid, and the staining-fluid (carbolised fuchsin solution) is used in the same manner as the *Ehrlich-Weigert* fluid. If the staining-fluid



FIG. 60.—Tubercle Bacilli from Sputum (eye-piece $\times 5$, objective $\times 15$, homogeneous immersion; open condenser).

be warmed, the specimen can be prepared in a few minutes. The surrounding tissues and remaining fungi (non-pathogenic) had best be stained with a watery solution of methylene-blue. The tubercle-bacilli then appear red, all other fungi and cells blue (fig. 61). It is a good plan to apply the staining fluid and acid solution by dropping them upon the cover glass held in forceps, rather than by immersing the preparation in a watch-glass.

[The Ziehl Neelsen method, with carbol-fuchsin, and decolorisation with 25 per cent. of sulphuric acid, is the simplest, and is now in general use in England. It may be carried out in the following way: (1.) A few cc. of carbol-fuchsin solution are heated to boiling and poured into a watch-glass. (2.) The cover-slip, prepared in the usual way, is floated for one to two minutes' preparation downwards on the hot solution. (3.) It is then removed and washed alternately in water and in a 25 per cent. solution of sulphuric acid until the film is colourless or

very faintly pink. (4.) The cover-glass is then floated for a half to one minute on Löffler's methylene-blue in a watch-glass, when it is removed. (5.) Washed, dried, and mounted for inspection.]

Koch-Günther Method.—For more than a year the author has exclusively employed the *Koch-Günther* method⁵⁷ and has found it reliable in every respect.

A mixture of 100 cc. of water and 4 cc. of pure aniline oil, thoroughly incorporated by agitation, passed through a wet filter, and mixed with 11 cc. of fuchsin solution.

The liquid must remain standing for twenty-four hours—a length of time which renders it advisable to keep some of the solution always ready—after which it is filtered and will then be fit for use.

The dry cover-slip preparations made ready in the usual manner are warmed in the solution until bubbles begin to form. Then, after a few



FIG. 61.—Tubercle-Bacilli, stained by the Ziehl-Neelsen method (eye-piece III., objective Zeiss $\frac{1}{2}$, homogeneous immersion, Abbe's mirror, open condenser).

minutes have elapsed, they are decolorised by immersion for a minute in a solution containing 97 cc. of 96 per cent. alcohol and 3 cc. of concentrated, chemically-pure hydrochloric acid, per 100 cc. The final staining is effected with an aqueous solution of methylene-blue.

Other methods which have been found useful clinically are those of Czaplewski, the Fränkel-Gabbett method (a modification of Günther's), and Biedert's.⁵⁸ The first has no advantage which does not belong also to the Ziehl-Neelsen method. The second is rapid and simple, but not entirely satisfactory in its results.

[Gabbett⁵⁹ employs two solutions:—(a.) *Magenta solution*: Carbolic acid (5 per cent.), 100 cc.; magenta, 1 gramme; spirit, 10 cc. Mix and preserve in a stoppered bottle. (b.) *Methylene-blue solution*: Sulphuric acid (25 per cent.), 100 cc.; methylene-blue, 2 grammes. The prepared cover-glasses are floated on a little of the magenta solution in a watch-glass. The latter is heated until the vapour can be seen rising from the surface. The cover-glasses are then removed,

washed, and placed for one or two minutes in the methylene-blue solution; after removal they are again thoroughly washed, dried, and mounted in xylol balsam. The whole process can be accomplished in a very few minutes.]

Biedert's method is very serviceable where the sputum contains but a few bacilli. Ten to twenty cc. of the sputum are boiled in a watch-glass with a little dilute caustic soda; water is added and the mixture again boiled until it is thinly fluid. It is then allowed to stand in a conical glass for two or three days, after which the sediment is added to a small quantity of egg albumin, and examined for bacilli in the usual manner. The addition of an alkali renders the bacilli more difficult to stain, and the specimen must therefore be left for a longer time than usual in the fluid. The use of *Stenbeck's* sedimentator,⁶⁰ or of the centrifuge, facilitates the research.

*Kuhne's*⁶¹ method may be mentioned, but it has been affirmed that it is unsatisfactory.

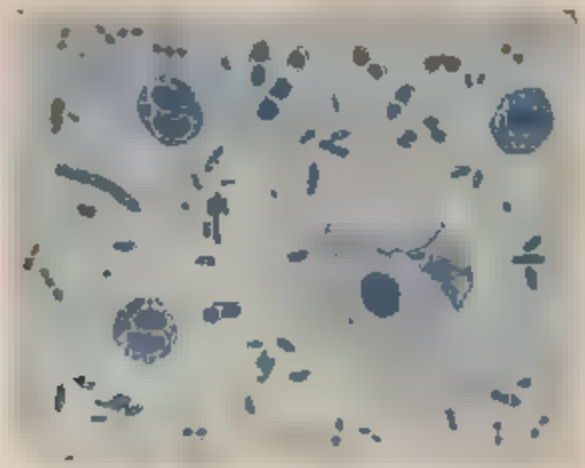


FIG. 62.—Pneumococci (eye-piece III, objective $\frac{1}{2}$, *Reichert*, homogeneous immersion, *Abbe's* condenser, without diaphragm).

[*Gibbes'*⁶² double stain is much used in England. It dispenses with the necessity of washing the preparation in acid and can be rapidly applied. It is made thus: (a.) Rosanilin hydrochloride, 3 grammes, and methylene-blue, 2 grammes, are rubbed up together in a mortar; (b.) aniline oil, 5 cc., is dissolved in alcohol, 20 cc. (b.) is slowly added to (a.) in the mortar until all the stain is dissolved; distilled water, 20 cc., is then added slowly while the mixture is stirred. The stain is now ready, and may be kept in a well-stoppered bottle. To use it a thin layer of the sputum is obtained on a cover-glass in the usual way. A little of the mixture is warmed in a watch-glass, and the cover-slip—sputum surface downwards—is floated on the warm staining-fluid for four or five minutes. It is removed and washed with spirit by means of a wash bottle, until the latter ceases to remove any of the dye. The preparation is allowed to dry, and mounted in xylol balsam.

Devices for estimating the number of bacilli present in a specimen of the sputum are both impracticable and devoid of clinical interest.

It is sometimes useful to be able to cultivate the bacilli directly from

the infected sputum. To do this *Kitasato*⁶³ advises that the expectoration raised by a deep cough be received directly into a sterilised Petri saucer, a portion of this detached on a probe washed in some 10 cc. of water, teased out with the usual precautions, and cultivated on blood serum or glycerine-agar. The resulting cultivations are round in contour, white, and raised above the surface of the medium—differing in appearance from those derived from the cadaver—these being in the form of small, scaly overgrowths.

We shall have occasion by-and-by to point out the great diagnostic significance which attaches to the presence of tubercle bacilli in the sputum.

2. Micro-organisms of Pneumonia.—*Klebs*,⁶⁴ *Eberth*,⁶⁵ and *Koch*⁶⁶ have described various micro-organisms, presumably of a specific character, as occurring in the lungs, and to be found in the sputum of pneumonic patients.

*Friedländer*⁶⁷ has further investigated this subject, and has succeeded in cultivating and propagating these micro-organisms, but their nature is not yet fully understood. They occur in stained preparations in groups of two, three, or four. They vary in size, have usually a distinct envelope, and present the form sometimes of short thick rods (*Friedländer*), sometimes of diplococci (*A. Fränkel*).

For the detection of pneumonococci (see upper part of fig. 62), *Friedländer*⁶⁸ recommends a method of staining very similar to that of *Günther* for the detection of spirilla in the blood (p. 52). The cover-glass preparation, made ready as described above, is passed three times through the flame of a Bunsen's burner, placed for a minute or so in a 1 per cent. solution of acetic acid, the fluid removed by blowing upon it through a pointed glass tube, the preparation dried in the air, and placed for a few seconds in a saturated aniline-water and gentian-violet solution, finally washed with water, and mounted on a slide for examination. When this is done a number of capsulated rod-like micrococci may be seen with the microscope.

From a series of experiments made by *Dr. Richter* it would seem that this method also answers remarkably well for the detection of fungi in exudations or transudations.

Gram's method (p. 47) will also serve to show the micro-organisms, which then appear chiefly as small diplococci (fig. 62, round the edges, and fig. 65), identical in appearance with the bodies supposed by *A. Fränkel* and *Weirselbaum*⁶⁹ to be characteristic of pneumonia.

We shall refer later to the pathological significance of these micro-organisms.

[3. The diphtheria-bacillus was found by *Kanthack* and *Stephens*⁷⁰

in the lungs in a large proportion of severe and fatal cases of the disease. The lungs were in many instances the seat of broncho-pneumonia. These observers doubt not that the bacillus is conveyed to the lungs, as it is to spleen and liver, by the blood, and that this event is not uncommon in cases of severe type.]

4. **Influenza-Bacillus.**—A specific micro-organism was found by *R. Pfeiffer*⁷¹ in the bronchial mucus of influenza patients, and cultivated by him. *Kitasato*⁷² obtained five generations of it by cultivating on glycerine-agar; and *Canon*⁷³ discovered similar fungi in the blood of persons affected, and by cultivation methods proved their identity with the bacilli of *Pfeiffer* and *Kitasato*.

According to *Pfeiffer*, the bacillus is of a thickness rather less than that of mouse septicæmia. It stains readily in Loeffler's methylene-blue solution with the aid of heat (p. 47) and with diluted Ziehl-Neelsen fluid, but not by *Gram's* method.

The bacilli are in prodigious quantity in the sputum (fig. 66), where they are usually lying free. In a later period of the disease they are found also in the contents of the pus-cells.

To cultivate the influenza-bacillus, a portion of sputum is taken, as described on p. 127, and implanted on glycerine-agar, or gelatine mixed with blood serum. On obliquely solidified glycerine-agar the cultures may be seen with a lens, after the lapse of twenty-four hours, like drops of water. They do not coalesce, but remain apart, and the bacilli develop freely. As a cultivation medium the author has used gelatine, to which, when fluid, a little blood has been added. *Nastjukow*⁷⁴ speaks well of egg-yolk-agar and egg-yolk-bouillon.

For the detection of the bacillus in the blood *Canon* proceeds as follows: A cover-glass preparation is made from the suspected blood, put to stain in *Chenzinsky's* eosin-methylene-blue fluid (see p. 67) for from three to six hours at 37° C., and subsequently treated with absolute alcohol for five minutes or more.⁷⁵

5. **Actinomyces.**—This fungus settles occasionally in the lungs, and may then be apparent in the sputum. *Baumgarten*,⁷⁶ *Israel*,⁷⁷ *Jekinowitsch*,⁷⁸ *Kuscher*,⁷⁹ and *Paltauf*,⁸⁰ [and *Lindt*] have described such cases. *Paltauf* has found the characteristic granular masses in the sputum in cases of actinomycosis, and their peculiar ray-like arrangement is made evident by the application of *Gram's* method. *Jekinowitsch* and *Kuscher* have shown that in the sputum the characteristic actinomyces occurs. For a description of the ray-fungus see the chapter on *Pus*.

6. **Plague-Bacillus.**—During the occurrence of epidemics of the plague the examination of the sputum for plague-bacilli will become important, since—as was shown in particular by the observations of the German Plague Commission⁸¹—this infectious disease may also assume

the characteristics of pneumonia, in which event the sputa of the patient will contain plague-bacilli in abundance (see Chapter I.).

It should be mentioned here that organised bodies have often been observed of late years in the sputum of whooping-cough, some of which may have a causative connection with the disease. Thus *Deichler*⁸² has found certain amoeboid cells (protozoa) in such sputa, but his statement needs confirmation. *Burger* and *Letzerich* established the presence of bacilli, and *Afanassiew*⁸³ found in the sputum of affected children bacilli to which he attributed an exciting influence; and since his opinion is supported by *Smtschenko*⁸⁴ (who performed careful cultivation experiments to elucidate the subject), it is probable that the bacillus of *Afanassiew* has really a close relation to whooping-cough. It should be mentioned, however, that *Cohn* and *Neumann*⁸⁵ came to quite a different conclusion. They failed to find the bacillus, or found it so seldom, that its connection with the malady was not apparent.

2. **Infusoria.**—Infusoria were first found by *Kannenberg*⁸⁶ in sputa derived from pulmonary gangrene. They were of the monad and cerco-monad varieties. They occurred usually enclosed in small yellow droplets, which also held margarine crystals, and exhibited sluggish movements. Such organisms are described at length in the chapter on *Fæces*.

For their detection in the sputum *Kannenberg* has adopted the following method:—A little of the fungoid substance mentioned above is placed beneath a cover-glass on a slide, and spread out by pressure in a very thin layer. A few drops of a 1 per cent. solution of common salt are then added. The cover-glass, with some of the preparation on its surface, is next dried, and afterwards put to stain in a watery solution of methyl-violet, washed in water, and, while still wet, placed in a concentrated solution of acetate of potash. By this process the protoplasm of the monads is stained a beautiful blue.

3. **Vermes.**—It very seldom happens that ascarides are found in the sputum, and cysts of echinococcus in a perfect state have been but rarely known to be expectorated. *Eichhorst*⁸⁷ and also *Hochsinger*⁸⁸ have described cases in which this happened. Diagnosis in such cases cannot be attended with difficulty. The cyst, however, is generally discharged in fragments. These appear to the naked eye as whitish-yellow shreds, and may be recognised microscopically by their uniformly striped texture (fig. 63). The discovery of the hooklets of echinococcus in the sputum is a fact of great importance in diagnosis. They may be readily recognised by their characteristic form (fig. 63). Charcot-Leyden crystals in great quantity commonly accompany them.

It would appear, too, that the expectoration may contain the eggs of *Distoma hæmatobium*. The author is indebted to Dr. Schiess Bey of Alexandria for a specimen, which plainly shows that this parasite

settles in the lungs, and it follows that when the pulmonary structure breaks down it may be expelled with the sputum. Similar observations had already been made by *Manson*.⁸⁰

A special interest attaches to the *Distoma Westermanii* or *pulmonale*.⁸⁰ The worm is cylindrical, of a reddish-brown colour, with a prickly surface, 8-10 mm. long and 4-6 mm. broad. The genital pore is situated behind the ventral sucker. The intestinal processes are unbranched. The uterus is globular, and lies immediately behind the genital pore. The male organ is bipartite, and placed in the hinder part of the body. The parasite infests the lungs of cats and the predatory feline tribe, but is found also in those of men, and, in particular, amongst the Japanese. It is then a cause of haemoptysis, and the eggs are seen in the sputum. They are brown, 0.08-0.1 mm. long and 0.05 mm. broad, and have blunt ends.

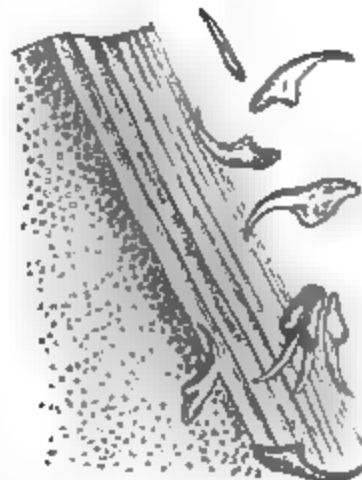


FIG. 63.—*Echinococcus* Hooklets and Membrane of Hydatid Cyst (eye-piece III., objective 8*a*, *Reichert*).

10. Crystals.—Very many forms of crystals have been found in the sputum, but for the most part their discovery is without great significance in diagnosis.

1. Charcot-Leyden Crystals.—We shall consider these bodies first, for the reason that they seem to possess some pathological interest.

*Leyden*⁹¹ often found crystals in the sputum of asthmatic patients. They abounded chiefly in the semi-solid greyish-yellow pellets discharged during a paroxysm. The crystals were colourless and of the pointed octahedral form. They were insoluble in cold water, æther, alcohol, and chloroform, but dissolved readily in acetic and mineral acids, alkalies, warm water, and ammonia. They are identical with the crystals to be seen in *post-mortem* blood, and described at p. 120, and with those of the semen, and with others that are met with in the faeces in cases of ancylostomiasis. *Schreiner*⁹² believes that such crystals are the phosphate of a new base, which, according to the researches of

Ladenburg and *Abel*,⁹³ is probably identical with *æthylenimin*, or *diæthylendiamin*,⁹⁴ though this identity is disputed by *Th. Kohn*⁹⁵ on very good grounds.

Leyden believed that the crystals were the exciting cause of the asthmatic paroxysm. *Friedreich*⁹⁶ and *Zenker* found them in expectorated fibrin-coagula, and *Bizzozero*⁹⁷ in the sputum of acute bronchitis from patients who were not subject to asthma. The author can corroborate the statement of these observers. [Troup⁹⁸ asserts also that neither spirals nor Charcot-Leyden crystals are pathognomonic of any one disease, but points out that the former are invariably to be found in asthmatic sputa, and by their presence cause the paroxysms. A characteristic feature of the conditions in which both spirals and crystals are found is marked desquamation of the mucous surface of the bronchioles, and the same author refers the origin of Charcot-Leyden crystals to the altered cells which are thus stripped off.]

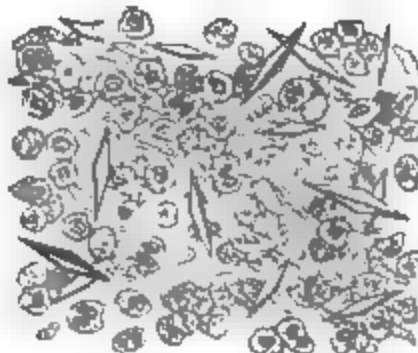


FIG. 64.—Charcot-Leyden Crystals, from the Sputum of an Asthmatic Patient
(eye-piece III., objective VII., Hartnack).

2. Hæmatoidin Crystals.—Crystals of hæmatoidin, described by *Virchow*, *Friedreich*, and *Schultze*,⁹⁹ occur in the sputum as ruby-red rhombic prisms, either solitary or in groups, or as needles or clusters of needles. These crystals, or fragments of them, are often found enclosed within the substance of white blood-corpuscles (fig. 53, e). Under such circumstances hæmatoidin is also seen, either free or in the interior of the white blood-corpuscles, as a mass of pigment in which no trace of the crystalline formation can be discovered.

When hæmatoidin crystals appear in the expectoration, we may conclude that blood has been effused and suffered to remain for some time in the air-passages, or that an abscess has perforated the lung. They occur most abundantly, therefore, in phthisis after hæmoptysis, when pulmonary clots are in process of absorption, very often in pulmonary abscess, and when an abscess or suppurating hydatid cyst has discharged into the lung. When the crystals are contained in cells, they point to a previous hemorrhage; but when free hæmatoidin is

present in considerable quantity, the inference is that an abscess has discharged from some neighbouring organ in the lung.

3. Cholesterin Crystals.—Crystals of cholesterin were found in the sputum of tuberculosis by *Biermer*,¹⁰⁰ and in pulmonary abscess by *Leyden*.¹⁰¹ In connection with the former disease they are often to be seen there, but in very small quantity. The author has observed them in great numbers in a girl who had a hydatid abscess in the lung, and in a man suffering from chronic pulmonary inflammation. According to *Black*,¹⁰² cholesterin crystals are plentifully present in the remains of old inflammatory exudation.

These crystals are distinguished by their powerful refractive action on light. They are in the form of large, and often irregular, rhombic tables, which have a tendency to cohere in groups. They are readily soluble in æther; insoluble in water, acids, and alkalies (fig. 130). When the crystals are treated with dilute sulphuric acid and a little tincture of iodine, a violet colour is formed, which presently changes to blue, green, and red. With sulphuric acid alone, they become first yellow and then green, these colours spreading from the edges.

Pathologically, these bodies are of little consequence. They are perhaps most apt to form when pus from some other organ has burrowed into the lung, and lodged there for a time before being discharged through the bronchi.

4. Fatty Crystals (Margarine Needles).—These are seen chiefly in putrid bronchitis and pulmonary gangrene. They characterise the discharge of unhealthy pus within the lungs, and belong to bronchiectasis and occasionally to tubercle. They occur either singly or in clusters as long sharp-pointed needles, and are occasionally vaulted or saddle-shaped. They dissolve readily in æther and boiling alcohol, and are insoluble in water and acids. This character facilitates their recognition (fig. of Semen, Chapter IX.). Recently the author has found them in plugs, which evidently originated in the crypts of the tonsils (see p. 110).

It follows from their occurring in so many different diseases that the discovery of these crystals in the sputum lends but little aid to diagnosis.

As to their chemical constitution, they consist, most probably, of the sodium, potassium, calcium, and magnesium salts of the higher fatty acids, as palmitic, stearic, &c.

5. Tyrosin Crystals.—*Leyden*¹⁰³ first found tyrosin crystals in the sputum of a girl with putrid bronchitis, and again in the case of a man who had an empyema discharging into the lung. They appear microscopically as fine needles scattered separately or in clusters. As a rule, they occur sparingly in fresh sputum, but form in greater quantity when the specimen has been allowed to stand for some time.

Leyden and *Kannenberg* are of opinion that a profusion of tyrosin crystals implies the existence of a perforating abscess of the lung.

It should be observed, however, that in many instances substances which have been taken for tyrosin were, in reality, lime and magnesia salts of the higher fatty acids.

Tyrosin is usually associated with leucin in the sputum. The latter occurs in faintly lustrous spherical particles (*R. Fischer*).¹⁰⁴ It has the same import as tyrosin.

For the detection of both these substances the method described in connection with the *Urine* may be employed.

6. Oxalate of Lime.—*Fürbringer*¹⁰⁵ has reported a case of diabetes in which the sputum contained large quantities of oxalate of lime. It occurs either in the shape of octahedral (envelope-shaped) crystals (fig. 118), or as an amorphous conglomerate. *Ungar*¹⁰⁶ found this body in the expectoration of a knife-grinder, aged twenty-eight, who had suffered for years from asthma.

The crystals of oxalate of lime are readily known from the fact that they are soluble in mineral acids, and insoluble in water, alkalies, organic acids, alcohol, and æther.

7. Triple Phosphate.—The characteristic coffin-lid crystals have occasionally been met with (fig. 119). They are soluble in acids of all kinds, and are, therefore, only to be found in the alkaline sputum. They are for the most part a product of the decomposition of proteids, attended with the liberation of ammonia. They are commonly to be seen in purulent exudation, and consequently are very plentiful in the expectoration from a discharging abscess.

Other crystals are also occasionally observed in the sputum. Fig. 53, d, represents some that were expectorated in a case of phthisis. They gave the chemical reactions of carbonates (carbonate of lime), the evolution of gas on the addition of acids, &c.

III. CHEMICAL EXAMINATION.—The chemical examination of the sputum throws little light upon disease.

1. Proteids and Allied Substances.—Of these, serum-albumin and large quantities of mucin and nuclein (*H. Kossel*)¹⁰⁷ are normally present. Peptone occurs in the expectoration of pneumonia and other purulent conditions—in general, in all cases where the sputa contain pus-cells in abundance.¹⁰⁸

The best method for the detection of proteids is that recommended by *Hoppe-Seyler*¹⁰⁹ as a test for albumin in serous fluids.

The “prune-juice” expectoration of pulmonary œdema is very rich in serum-albumin. For the detection of this body, the sputum may be

extracted with very dilute acetic acid, and the filtrate tested with ferrocyanide of potassium, when its presence will be shown by turbidity or a precipitate.

To estimate the quantity of albumin *R. v. Jaksch* and *F. Lanz*¹¹⁰ proceed thus: A known weight of sputum is placed in the v. Jaksch flask and submitted to *Kjeldahl's* process. The quantity of nitrogen obtained is measured, and the figure expressing this is multiplied by 6.25. The product is the amount of albumin present; but since the constituent proteids are very various, an estimate based upon the quantity of nitrogen is only approximately accurate. The results which the author arrived at in this way were very inconstant. Thus in tuberculosis, the mean of twenty cases, with thirty-six estimations, showed 0.6795 grm. of nitrogen in 100, while in pneumonia the proportion was greater, reaching as high as 1.7784 per cent. The first figure corresponds to 4.2468 per cent., the second to 11.1150 per cent. of albumin. Equally varying results were obtained by *Starkow* and *Fr. Müller*.¹¹¹ Further researches by *Lanz* show that in pulmonary tuberculosis there is a great loss of proteids from the body.

2. Volatile Fatty Acids.—Various volatile fatty acids may occur in the sputum, and, notably in gangrene of the lung, acetic, butyric, and sometimes caproic acids are met with.¹¹² In testing for these bodies, the sputum is diluted with water and phosphoric acid added. The volatile constituents are driven off by distilling them in a vessel heated by steam. The distillate is then tested in the manner laid down in the chapter on *Fæces*. Some fat can be obtained from all sputa, and those of tubercle patients contain a large quantity.¹¹³

To separate the fats and non-volatile fatty acids, the sputum is acidulated and extracted with æther, and by repeatedly shaking up the æthereal extract with a watery solution of sodic carbonate, the acids are converted into their corresponding salts, which remain dissolved in the watery solution. The æther is pipetted or siphoned off, and the fats obtained after evaporation of the æther. The sputum of pulmonary gangrene contains several members of the aromatic group, as indol, phenol, and scatol.¹¹⁴

3. Glycogen.—This body has repeatedly been detected in the sputum by *Salomon*.¹¹⁵ Its presence may be demonstrated by *Brürke's* method.

4. Ferment.—*Filehne*, *Stolnikow*, and *Stadelmann*¹¹⁶ found that the sputa, especially in pulmonary gangrene and putrid bronchitis, contain a ferment resembling in its action one of the pancreatic ferments. It is soluble in glycerine, and may be extracted by that body from the sputum. *Escherich*¹¹⁷ has been able to demonstrate its presence in the sputum in all cases of extensive destruction of lung-tissue.

5. Inorganic Constituents.¹¹⁸—These are chiefly—

1. Chlorides : of sodium and magnesium.
2. Phosphates : of soda, lime, and magnesia.
3. Sulphates : of soda and lime.
4. Carbonates : of soda, lime, and magnesia.
5. Per-salts of iron (rarely)—phosphate of iron.
6. Silicates.

The inorganic matter of the sputum has little bearing on clinical diagnosis. For its analysis the organic constituents must first be removed by incineration, when the usual tests may be applied to the ash. (For further information consult *Hoppe-Seyler* and *Thierfelder's Handbuch der physiologisch- und pathologisch-chemischen Analyse*, 6th ed., p. 304.)

IV. THE CHARACTER AND CONSTITUTION OF THE SPUTUM IN THE PRINCIPAL DISEASES OF THE LUNGS AND BRONCHI.**1. Diseases of the Bronchi.**

1. **Acute Bronchitis.**—At the outset the expectoration is viscid, white, and scanty, and occasionally streaked with blood. Microscopically it exhibits very few cells, *and is devoid of specific fungi (tubercle-bacillus, &c.).* At a later period it becomes more abundant, assumes a pale-green tint, and under the microscope is seen to consist chiefly or entirely of pus-cells. Elastic fibres are never present.

2. **Chronic Bronchitis and Bronchiectasis.**—The expectoration, which is copious, green in colour, and without any characteristic odour, is formed microscopically almost entirely of pus-cells, with which are mixed a considerable number of fat-holding epithelial cells and myelin particles, and a host of non-pathogenic micro-organisms. When ulceration of the bronchi takes place in the course of chronic bronchitis, leading to bronchiectasis, a very abundant expectoration is discharged in the morning. The fluid, which is usually thin, tends to arrange itself in three layers when allowed to settle. Of these, the topmost is froth, the next consists of watery fluid, and the third, which is of greater consistency, is formed almost exclusively of cells.

In chronic bronchitis complicated with asthma, the sputum during the paroxysms, and immediately afterwards, exhibits spirals (see p. 119), and *Charcot-Leyden* (p. 113) and other crystals.

3. **Fœtid Bronchitis.**—The expectoration of this disease is thick, greenish-brown, and has a very disagreeable sweetish odour. Microscopically it contains a profusion of micro-organisms of various kinds,

and very often contains large tufts of fungi, which colour blue with iodine and iodide of potassium solution, epithelium—usually in the process of fatty degeneration—and fungoid plugs (p. 125), but no elastic fibres, no shreds of alveolar tissue, and no specific fungi.

*Lumniczer*¹¹⁹ has separated by means of *Koch's* plate cultivations (see Chapter X.) a number of micro-organisms from the sputum, amongst them *Staphylococcus pyogenes citreus* and *albus*, *cereus* *flavus* and *albus*, and *diplococci*; and also a fungus, the colonies of which, when developed in nutrient agar-agar, emitted the odour belonging to the expectoration of putrid bronchitis. It is a spore-forming bacillus, $1.5-2 \mu$ in length, thickest in the middle, and rounded off at the ends. It was also seen ready formed in the sputum. When introduced within the lungs and bronchi of rabbits it excited inflammation there.

Læbisch and *Rokitansky*,¹²⁰ by means of *Baumann* and *Udransky's* benzol test, have detected cadaverin (pentamethylenediamin) and another diamin in these sputa.

4. Plastic Bronchitis.—The sputum contains fragments of the croupous membrane and coagula of fibrin, with which are entangled a profusion of epithelial cells and fungi (fig. 58). From these appearances, and in the absence of pneumonic symptoms, the condition is readily diagnosed. Occasionally it has to be distinguished from chronic fibrinous bronchitis. The distinction will be made on clinical grounds.

2. Diseases of the Lung Tissue.

1. Pulmonary Tuberculosis.

(a.) **Miliary Tuberculosis of the Lung.**—The sputum in this condition is merely that of acute catarrh. *Tubercle-bacilli* are never to be found in it.

(b.) **Acute Tubercular Infiltration of the Lung.**—An early diagnosis of the condition is of the utmost consequence, and happily it has been rendered possible by *Koch's* epoch-making discovery of the tubercle-bacillus. Two forms of the disease may be distinguished, according as the symptoms partake of the typhoid or of the pneumonic character.

(a.) **The Typhoid Type.**—The symptoms are rigors at the outset, persistent high temperature, enlarged spleen, extensive roseolar rash, which at times suggests that of typhus exanthematicus, and commonly profuse diarrhoea. The physical signs are those of severe catarrh in both apices. There is no dyspnœa. The pulse is frequent, respiration is but slightly accelerated, and lividity is not marked. The sputum is viscid and scanty. It contains but little tissue-débris, and only a few bacilli, which, moreover, are always provided with spores.

After the lapse of a few days, dulness on percussion and bronchial breathing may be obtained over the apices. The sputum becomes purulent, and when again examined is found to contain *swarms of tubercle-*

bacilli. There are also to be seen elastic fibres, showing an alveolar arrangement, and a large quantity of epithelium. The physical signs now give evidence of one or more large cavities, and the fever assumes a remittent character. Death generally ensues in three or four weeks, the mode of dying being that of chronic tuberculosis.

(β.) *The Pneumonic Type* begins with sustained high temperature, marked lividity, and accelerated breathing. The physical signs point to catarrh of both apices, and the sputum contains a few bacilli. Later, the characteristic symptoms of infiltration of the lungs set in; the expectoration becomes more profuse, and the contained bacilli more numerous. The disease runs a very rapid course, often lasting only for some days. Anatomically, the condition is one of acute tubercular infiltration of both lungs. That, as a matter of fact, the sputum, in cases of this form of tuberculosis, may exhibit all the characteristics of that of croupous pneumonia, is proved by a case, from the author's clinic, published by *Hoke*.¹²¹

(c.) **Chronic Pulmonary Tuberculosis.**—The diagnosis of this disease has been greatly facilitated by Koch's discovery of the bacillus of tubercle. As a result of his experience of many hundred cases in the course of late years, the author has no hesitation in affirming the principle so often laid down by other authorities, that *the presence of the tubercle-bacillus in the sputum invariably implies the existence of tuberculosis*. Let this fact be granted, and no more is needed at once to emphasise the importance of that recent contribution to our knowledge, and to impress upon the physician the necessity of rendering himself familiar with the very simple methods by which the bacillus may be identified (p. 126). Experience has already accumulated certain data. In the first place, it must be mentioned that in most, though not in all, cases the abundance of the bacilli is in proportion to the severity of the disease; and again, the proliferation of bacilli is apt to run parallel to the intensity of the febrile symptoms. The supervention of haemoptysis is attended with an apparent (*H. v. Frisch*¹²²) diminution in their number, but this may well be owing to the fact that the sputa are in such a case diluted by the profuse admixture of blood.

It is only when the tubercular process runs the very rapid course described above that the bacilli, then usually carrying spores, are so abundant as to fill the entire field.

It was held at one time that the appearance of elastic fibres in the sputum was a sure token of incipient tuberculosis, but more recent research has shown that they may be found equally in the course of pulmonary ulceration of whatever kind; and so with other manifestations to which great importance was formerly attached. The signifi-

cance of all or many of them is very slight in comparison with that belonging to the specific micro-organism, upon the detection of which alone a positive diagnosis can be formed. [By the method of examining the sputum in sections already referred to (p. 115), Gabritschewsky¹²³ has observed giant-cells in three out of four cases of pulmonary tuberculosis.]

It does not necessarily follow from the presence of the bacillus in the sputum that a fatal termination is impending. The author has met with such cases in which the patient eventually recovered, but they are very rare. And doubtless an atmosphere crowded with the bacilli of tubercle (as that of the consumptive ward of a hospital) must necessarily give but little chance of cure. Possibly this is the reason why the hospital physician has so seldom an opportunity of observing it.¹²⁴

Allusions may be made here to Koch's process¹²⁵ for the detection during life of tubercular proliferation in inaccessible parts. The diagnostic value of the discovery is not yet finally settled. Numerous clinical observations have shown that persons affected with tubercle-bacilli do in fact exhibit a reaction when injection has been made upon them, while the healthy, and those subject to other diseases, generally fail to do so. The process, however, is *neither entirely trustworthy nor altogether free from danger*, and therefore cannot be recommended for purposes of diagnosis in the human subject. In the case of animals (cattle), however, it appears to be reliable.

2. Chronic (Non-Tubercular) Inflammation of the Lung.—Under this heading may be grouped all those conditions which would formerly have been thought to present the typical character of tuberculosis—fever, night-sweats, &c.—but in which repeated search will fail to detect the bacillus in the sputum. In one such case the autopsy showed caseating masses of considerable extent, but these, even to the naked eye, were sufficiently discernible from tubercular deposit.

As regards the sputum, its essential characteristic is the *absence of tubercle-bacilli*. It further contains a large quantity of elastic tissue and a profusion of epithelium cells, and especially of their myeloid derivatives. If we may infer from the very slender data which have as yet accumulated, the disease is attended throughout with slight fever, and death by asthenia occurs sooner or later.¹²⁶ The author is convinced that, with a little more care, we should learn to recognise the non-bacillary form of phthisis in many instances where at present it is not thought of.

3. Croupous Pneumonia.—At the outset of this disease the expectoration is very scanty, of a white colour, and here and there streaked with blood. Microscopically, it exhibits only a small number of red

and white blood-corpuscles ; it usually, though not always, contains the specific micrococci which will presently be described.

Later, and in some cases only a few hours after the initial rigor, the sputum has a rusty tinge. It is usually also at this period remarkably tenacious, and adheres firmly to the spit-box. When placed under the microscope, it is seen to contain but a small number of much-shrivelled blood-corpuscles, the red ones arranged for the most part in rows. They are too few to account for the colour of the sputum, which is due rather, as *Traube* surmised, to the presence of dissolved haemoglobin. In addition to these there are some leucocytes, and the alveolar epithelium which has been noticed in another place (p. 117). In rare cases the spiral bodies already described and shreds of fibrinous coagula may also be observed.

The colour of the sputum at this stage, and even later, may be grass-green, even though there be no jaundice. Professor Nothnagel¹²⁷ is of opinion that under such circumstances the colouring matter of the blood is changed into bile pigment. At his desire the author has examined some of these grass-green sputa. They were dissolved in a mixture of alcohol with a little chloroform, which was then filtered and the filtrate evaporated, when a substance remained behind which presented all the characters of biliverdin. We may assume, therefore, that the green colour of the sputum in such cases is due to the conversion of haemoglobin or haematin into bilirubin, a change which is the less remarkable if we consider the very close chemical relations which these two bodies bear to one another. The bilirubin formed would then be oxidised to biliverdin in the lungs.

Traube maintains that the grass-green sputa belong especially to subacute pneumonia, and that they also occur when a pulmonary abscess forms in the course of pneumonia.

It would appear from the researches of *Rosenbach*¹²⁸ in Nothnagel's clinic at Jena, that the sputa may be tinged green by certain micro-organisms, most probably the *Micrococcus chlorinus*,¹²⁹ altogether independently of pneumonia. This happens under many circumstances ; and the fact has in itself no clinical significance.

Later in the course of pneumonia the expectoration becomes more abundant and watery. Its colour changes from brownish-red to a saffron- or citron-yellow, the alteration in most cases depending upon a change in the character of the blood pigment. Fibrinous coagula, and occasionally also spirals, now become more numerous.

It must be noted, however, here, that such saffron- or citron-yellow sputa do not belong exclusively to pneumonia. *Renz*¹³⁰ has recorded a case of tuberculosis, in which microscopical examination showed a profusion of haematoidin crystals in the expectoration, which was of a

yellow ochre tint. *Löwy*¹²¹ also has called attention to a form of sputum essentially distinct from that of pneumonia, though, like it, characterised by a yellow colour, which generally first appears after the sputum has been voided. It occurs, according to *Traube*, almost exclusively during the summer months in connection with tuberculosis, pleurisy, and pleuritic exudation. The colour in this case is due to the presence of micro-organisms, and no special importance attaches to the condition apart from its liability to be taken as a sign of pneumonia.

In the later stages of pneumonia, the fibrinous coagula disappear largely from the sputum, and the red corpuscles and leucocytes are greatly lessened in number, and show fatty degeneration. There may still be a few spirals, and elastic fibres are to be met with where ulceration is in progress. Finally, alveolar epithelium, either very fatty or hyaline (?) [*Feuerstock*¹²²], and commonly, too, exhibiting the myelin forms, is present in great abundance.

Lastly, if the disease tend towards recovery, the colour begins to disappear from the sputum, which also under the microscope exhibits a constantly diminishing proportion of epithelium (still, however, fatty), until at last a time arrives when it is indistinguishable from that of an ordinary bronchitis.

The subject under consideration would not be complete without some notice of the position which may be accorded to the pneumonia-coccus of *Friedländer*, when studied as an adjunct to diagnosis. We are not here concerned with the discussion of its effects as an agent of disease, and shall confine ourselves strictly to its significance as a symptom. Upon this point the author has collected the records of a large number of cases.

It should be stated in the first place that, in every case of pneumonia to which they have been applied, the methods of *Friedländer* and *Gram*, already described, have disclosed the presence in the sputum of bodies resembling the microbe of pneumonia. Since this is true also of cases of central pneumonia, which are always so difficult to diagnose at the outset, it becomes in the highest degree desirable to assign to these forms their proper significance. Pure cultivations of pneumonia-cocci are sometimes to be seen in the sputa of the disease (see fig. 65). It may, perhaps, be assumed that in doubtful cases their presence will confirm the suspicion of pneumonia; but inasmuch as *Friedländer's* microbes (or rather, it should be said, certain forms which closely resemble these) have been found in the expectoration of other conditions, such as chronic bronchitis and bronchiectasis, and in the buccal fluids and sputum of health, we are never warranted in basing a diagnosis on such manifestations alone. It may well be that the micro-organisms in question are not in all cases identical, and that by improved methods of

cultivation we shall be in a position to discriminate them; but at present we are unable to do so. The researches of *Pansini*¹²³ have shown that cultivation-results are not decisive, since fungi resembling the *Fränkel-Weichselbaum* coccus may be derived from sputa which are not pneumonic. Moreover, it would appear from the observations of *Fränkel*¹²⁴ and *Weichselbaum*¹²⁵ that there are several distinct microbes which are capable of giving rise to the symptoms of pneumonia. The researches of *Neumann*¹²⁶ have still further extended the limits of the variety which may be observed. It should, however, be mentioned that the researches of *Fränkel*¹²⁷ and *Weichselbaum*¹²⁸ leave no doubt that amongst the numerous micro-organisms which are present in croupous pneumonia, a diplococcus is most commonly to be found (*Micrococcus*



FIG. 65.—Microbe of Pneumonia from the Sputum, coloured after Gram's method (compensation eye-piece IV objective apochromatic immersion $\frac{1}{2}$, Zeiss).

of pneumonia, *A. Fränkel*; *Diplococcus pneumoniae*, *Weichselbaum*) (fig. 65).

Fränkel,¹²⁹ *Fou, Bordoni Uffreduzzi*,¹³⁰ and *Weichselbaum*¹³¹ have observed the same diplococcus in the purulent exudation of cerebro-spinal meningitis; but it would appear from the researches of *Weichselbaum* and *Goldschmidt*¹³² that there are also other forms which appear to be associated in a closer manner with this disease.

Lanz finds that the albumin in the sputum ranges from the maximum 1.7784 per cent. of N. = 11.150 per cent. of albumin, to the minimum 0.6753 per cent. of N. = 4.1206 per cent. of albumin; the mean of eleven observations being 0.9524 per cent. of N. = 5.8525 per cent. of albumin. When the crisis sets in the proportion sinks to 0.4 per cent. N. = 2.6 per cent. albumin, and even less.

To sum up our knowledge, it must be admitted that the subject of the pneumonia-microbe needs further elucidation. It is likely that there are several fungi which may excite the disease. The sputum of pneumonia occasionally exhibits pure cultivations of these fungi (fig. 65); but their presence must not be made the basis of a too confident inference. It is only under special circumstances that diagnostic interest attaches to any of them.¹⁴³

4. Influenza.—At the early stage of this malady the sputum has the ordinary microscopical characters of bronchial catarrh (p. 139). Later it is extraordinarily profuse (100–200 cc.), and consists of pure pus of a glutinous or viscous appearance, and in this the bacilli (fig. 66) may be detected. The author investigated the character and life-history of the bacillus, as opportunity occurred, during the epidemic which visited Prague at the end of 1893.¹⁴⁴ Using *Pfeiffer's* method, he found,



FIG. 66.—Influenza-Bacilli from the Sputum (*Zeiss's* eye-piece IV., objective apochromatic immersion 1.30).

microscopically, that bacilli were present in proportion corresponding to the severity of the clinical symptoms, the bacilli answering, in respect of appearance, number, and arrangement, to the description of the influenza-bacilli of that observer. He succeeded also, by cultivation, in producing colonies of similar micro-organisms; and, in addition to these, there were also broods of cocci.

The diagnostic significance of the bacillus, so far as it rests upon personal observation, may be deduced from the following facts. The sputum in twenty-two cases of tubercle, heart-failure, chronic bronchitis, and pneumonia *without* influenza, failed with *Pfeiffer's* method to exhibit the bacillus. Further, tubercular sputum examined by this method showed a remarkable scarcity of micro-organisms. In two cases, on the other hand, which very decidedly were *not* cases of influenza, large quantities of bacilli, resembling *Pfeiffer's* bacillus in appearance and dis-

tribution, were discovered. From this it follows that the bacillus is of diagnostic importance only when its cultivation-peculiarities have been elicited ; and it may be added that a sufficient clinical examination is the best and indispensable means at our disposal for diagnosing influenza. In the year 1895, when, taking the clinical symptoms as a guide, it would seem that but few cases of influenza came under his notice, the author did not in any instance observe the bacillus in the sputum.

5. Pulmonary Abscess.—The sputum of this condition generally looks like pure pus when seen with the microscope. It emits a faint and slightly foetid odour ; and it tends on standing to stratify in two layers, of which the lower consists of pus-cells, whilst the upper is watery and topped with froth. Microscopically the sputum is subject to variety, but in general it exhibits shreds of lung tissue, and very commonly elastic fibres with the alveolar disposition (fig. 54), fatty and disintegrated pus-cells, haematoxin—partly in the form of well-shaped crystals, partly as brown or reddish pigment-particles of varying size—often also cholesterin crystals ; these are equally abundant in cases where the pus has been of long accumulation. Tyrosin and balls of leucin are occasionally present ; fatty crystals are more commonly found ; and finally, a profusion of fungi of various forms, but these are never specific.

6. Gangrene of the Lung.—The sputum has a penetrating and extremely unpleasant smell ; is abundant, thin, and of a dull-green colour. It separates on standing into three well-marked layers. Of these, the upper is frothy, very turbid, and of a greenish-brown tint ; the middle thin, and of a watery or serous character ; whilst the lowest is opaque, viscid, and ranges in colour between brown and green. The last occasionally contains brown shreds of lung-tissue of varying size.

Microscopical examination shows that the upper layers contain but few formed elements. In the lowest layer, however, there is a large proportion of detritus, fatty globules of irregular size, rarely crystals, but most commonly haematoxin crystals and amorphous masses ; an enormous quantity of fungi, notably fission-fungi ; often large tufts of fungi, which stain blue with the iodo-potassic-iodide solution (*Leptothrix*), together with other forms, which resemble starch granules and stain similarly with the same reagent ; and sometimes also monads (*Kannenberg*).¹⁴⁵ It is important to notice the absence of elastic fibres. *Stadelmann*,¹⁴⁶ however, says they often occur in this kind of sputum. The sputum contains a ferment which acts like pancreatic juice, and it is this which probably causes the solution of elastic tissue (see p. 119).* *Bonome*¹⁴⁷ has constantly found in such cases the *Staphylococcus albus*

* [This view is opposed by Troup ("The Sputum," p. 48), who adduces important facts in support of his position.]

and aureus, which on this account he regards as the cause of this disease.

The researches of *Hirschler* and *Terray*¹⁴⁸ have disclosed a great variety of micro-organisms in this disease. They include a number of staphylococci—in addition to those already mentioned, *Staphylococcus pyogenes citreus*, and *cereus albus*—*Bacillus pyocyaneus*, and a micrococcus which thrives on gelatine, agar-agar, and blood-serum at 20°–24°, and on gelatine produces cultivations resembling four-leaved clover or a flower of six petals. It liquefies gelatine slowly, and on all nutrient substances develops a smell closely resembling that of gangrenous sputum. This fungus causes disease when inoculated upon animals. It assimilates aniline colouring matters of all kinds. It is stained with difficulty by *Gram's* method. It remains to learn whether it has any relation to pulmonary gangrene, and of what nature this may be. *Lanz's* researches show that this sputum is not very rich in albumin.

7. Pulmonary Oedema.—In this condition the sputum is abundant, thin, and watery, and according to the nature of the underlying process is either white and frothy (like soapy water), or of a rusty brown colour (like prune-juice). Microscopically it holds comparatively few cellular elements. The leucocytes and the few epithelial cells to be seen, especially in cases of acute oedema, are free from fatty changes. The red blood-corpuscles are few, and their number is inadequate to account for the deep colour of the expectoration. In one case the author believes that by digestion with water and subsequent filtration of the sputum he obtained the characteristic absorption-bands of methæmoglobin with the spectroscope. The presence of abundance of albumin may be shown by chemical means.¹⁴⁹

8. Hæmoptysis.—In cases of profuse hæmorrhage from the lung, the sputum consists entirely of bright-red frothy blood, with scarcely any trace of other tissues. When the hæmorrhage has diminished in intensity, the expectoration for several days continues to exhibit a reddish or reddish-brown colour. During this period leucocytes and epithelium are plentifully met with, the cells usually enclosing crystals and amorphous particles of hæmatoidin. The formation of small cavities in the course of tubercle is the commonest cause of pulmonary hæmorrhage. In addition to this we shall here notice only the rupture of an aneurysm into the bronchi, and remind the reader that a persistent hyperæmia of the lung may also give rise to this condition.

9. Hæmorrhagic Infarction.—In cases of recent hæmorrhagic infarction of the lung, single coin-shaped masses of blood, of a bright-red colour, and intimately mixed with froth, are expectorated. After the lapse of several days the sputa assume a brownish tint, and answer the description detailed in the last paragraph. Further, as a rule, they

contain a number of epithelial cells and leucocytes undergoing fatty degeneration. The cause of this condition is generally to be sought in functional or organic debility of the heart.

10. Pneumoconiosis.¹⁵⁰

(a.) **Anthracosis of the Lung.**—The sputum of tobacco-smokers and those who habitually breathe an atmosphere laden with soot always contains some proportion of carbon particles. In such cases the expectoration in the early morning is of a pearl-grey colour, and is brought up in pellets of viscid and remarkably tenacious substance. In a typical case of anthracosis the sputum ranges in colour from dark brown to black, and is somewhat abundant. Microscopically it exhibits particles of free carbon, readily discernible by their resistance to the action of acids and alkalies; and generally a quantity of leucocytes and alveolar epithelium, in each case choked with pigment particles. J. Wiesner¹⁵¹ has definitely proved that the black lung pigment consists of soot, so that there can no longer be any doubt that the black, amorphous particles present in the sputum are in most cases of this material.

(b.) **Siderosis Pulmonum.**—The sputum is usually brownish-black in colour, and has all the characters of that of chronic bronchitis. Under the microscope the contained leucocytes and alveolar epithelium are seen to be laden with a reddish pigment, which by its reaction with sulphide of ammonium (black coloration from formation of sulphide of iron), or with hydrochloric acid and ferrocyanide of potassium (prussian blue), may be known to consist of iron.¹⁵²

(c.) **Mason's Lung.**—Here, too, the sputum has generally the appearance of that of chronic bronchitis; but in addition it contains particles of dust, either free or enclosed within cells. Lime and gypsum dust are readily recognised by their chemical properties (see the chapters on *Faeces* and *Urine*), and ultramarine by its colour. In other cases evidence will be forthcoming from independent sources as to the nature of the substance in question.

CHAPTER V

THE GASTRIC JUICE AND VOMIT

I. EXAMINATION OF THE GASTRIC JUICE.—Like the saliva, the gastric juice is the product not of one but of several distinct sets of glands. It is composed of the fluid secreted by the pyloric glands of the stomach and that of the cardiac glands, these forming its active digestive ingredients ; and it includes also a portion of the buccal secretion which has been swallowed and is already partly changed in the process of digestion.¹

1. Naked-Eye Characters.—The gastric juice of man is a colourless fluid, generally clear, but it is occasionally turbid. Its reaction is acid.

2. Formed Elements.—Microscopical examination of the gastric juice at a time when the stomach contains little or no residue of undigested food shows single squamous epithelial cells derived from the upper part of the alimentary canal ; columnar epithelium very rarely and only in special instances ; fungi of various kinds, especially bacilli and micrococci ; and usually also yeast cells. *Abelous*² has distinguished sixteen, and *Lockhart Gillespie*³ twenty-four different micro-organisms. Amongst these are *Sarcina ventriculi*, *Bacillus pyocyaneus*, *Bacterium lactis aërogenes*, *Bacillus subtilis*, and others, all of which effect changes in the food—albumin, milk, or carbo-hydrates. From this fact it would appear probable that in the stomach, as in the intestines, certain of the fission-fungi are concerned in physiological functions. On the other hand, there can be no doubt that the gastric juice is the means, under appropriate conditions, by which a host of the most dangerous germs of disease are rendered harmless and destroyed. This property is derived from the free hydrochloric acid which hinders the development of such fungi.⁴ According to *Jaworski*,⁵ the microscopical characters of the fluid differ according as it has an acid reaction or otherwise. It should also be remarked that *Henschen*,⁶ in one case of carcinoma of the stomach, discovered infusoria (Monadines) in the contents of that organ.

If the examination be made while digestion is going on, the character of the gastric juice will approach that of which we shall have occasion

to speak later when dealing with the vomit, i.e., it will contain the remains of the food.

3. To Obtain the Gastric Juice.—*Leube*⁷ and *Külz*⁸ were the first to employ the gastric sound for this purpose in man. Its use is unattended with danger provided the tubes are elastic and too energetic aspiration be avoided. To procure the gastric juice for the purpose of chemical analysis, whether in health or disease, the following procedure has been recommended by *E. Schütz*.⁹

A time is chosen when the stomach is empty, so as to prevent, as far as possible, the admixture of impurities derived from the food, and to this end the morning is generally the best. A pliable gum-elastic sound, perforated at the end with a number of apertures not larger than a pin's head, and furnished with a lacquered handle, is introduced into the stomach, and pushed on until a slight resistance is encountered. A collar of horn is then slipped over its upper part, and grasped between the teeth of the person experimented upon, so as to keep the sound in position. After the lapse of about half a minute the handle is removed and the sound connected with a stomach-pump. The piston is now drawn out, and the projecting extremity of the tube being grasped with the fingers, the sound is withdrawn, and its contents discharged by the movement of the piston into a glass vessel. As already said, the danger of the operation is very slight when performed in this way; but it may be entirely obviated by introducing a mercurial manometer between the sound and the pump, and estimating beforehand the amount of pressure (as indicated by the position of the mercury) which may be employed without applying any considerable suction force to the mucous membrane of the stomach. *Gross*¹⁰ has devised a very serviceable modification of the stomach-tube.

It is advisable in many cases, and especially where children are concerned, to adopt the simpler method of *Boas* and *Eicard*,¹¹ which consists in applying pressure over the abdomen after the introduction of the elastic tube. By this means the fluid is forced into the latter, and may be drawn off as by a syphon.¹²

As a substitute for these methods, *Edinger*¹³ causes the patient to swallow pieces of sponge compressed and coated with gelatine, and retained by a thread which is held in the hand. *Späth*¹⁴ similarly uses scraps of elder pith, stained with appropriate reagents, or grains of shot having attached to them threads soaked in the reagent. *Bocci*¹⁵ has devised a little instrument by means of which 0.1 grm. of gastric juice may be withdrawn. *Sahli* and *Günsburg*¹⁶ employ tablets containing potassium iodide in a thin coating of gum, and connected by threads of fibrin. These are given to the patient, and the appearance of iodide in the saliva is taken to indicate the rapidity with which fibrin is absorbed.

4. Chemical Constituents of the Gastric Juice.—Of these, the most important are—(1.) Pepsin; (2.) Rennet, or milk-curdling

ferment; (3.) Inorganic and organic acids. Each of these is liable to undergo pathological change, both as to quantity and to quality. Those which affect the *pepsin* and *acid* constituents are of chief consequence in disease.

1. Pepsin.

(a.) *Detection of Pepsin in the Gastric Juice.*—As a test for pepsin, its property of changing proteids, e.g., fibrin, into peptone is turned to account. The best method of procedure is as follows:—10–20 cc. of the acid fluid, obtained as above, is taken, diluted with water, and filtered. To the clear filtrate a small quantity of well-washed blood fibrin is added, and the whole kept at a temperature of 40° C. If pepsin be present, the fibrin will be dissolved after a few hours. If, after the lapse of 10–12 hours, no change is apparent, or if an odour of putridity be given off by the fluid, it may be assumed that the latter is free from pepsin. Should it happen also that the secretion obtained from the stomach has an alkaline or but feebly acid reaction, it will be necessary, before applying digestion tests, to add to it its own volume of a dilute solution of hydrochloric acid (8 cc. of the fuming acid in 992 cc. of water).

(b.) *Quantitative Estimation of Pepsin.*—*Schütz'* method may be employed for the quantitative estimation of pepsin. It is founded upon a principle first enunciated by *Huppert* and *Schütz*,¹⁷ one of fundamental importance for the theory of digestion—namely, that under certain conditions at the disposal of the observer, the quantity of peptone formed is exactly proportional to the square root of the quantity of pepsin used. *Schütz* has taken as the pepsin-unit that quantity of the ferment which will yield 1 grm. of peptone under the conditions of his experiment, and expresses his results in terms of this unit. For further details the reader is referred to the original communication. A process recently advocated by *Hammerschlag*¹⁸ gives at best but approximate results, and is on this account not to be recommended for scientific purposes.

2. Milk-Curdling Ferment.—This ferment was first investigated by *Hammarsten*. It may be detected by the following process:—2–10 cc. of cow's-milk, of neutral reaction, is well boiled, and to it is added an equal quantity of gastric juice which has been carefully neutralised and filtered. The mixture is placed in a warm chamber, or on a water-bath heated to 30°–40° C. If the milk-curdling ferment be present, the casein of the milk will be precipitated in flakes after the lapse of 20–30 minutes. The ferment was found by *Schumburg*¹⁹ and *Boas*²⁰ to be invariably present in health, and absent in serious disorders of the stomach, as cancer and atrophy of the mucous lining. It is wanting in infants of from one to two days old; but *Raudnitz*²¹ established its presence in older children who were reared on cow's-milk. It was

regularly found in a series of investigations conducted on such subjects (*v. Jaksch*).

The researches of *Johnson*,²² *Boas*,²³ *Klemperer*,²⁴ *C. Rosenthal*,²⁵ *A. Johannessen*,²⁶ and *O. Sandberg*,²⁷ have thrown much light on the subject of the rennet constituent of the gastric juice. These tend to the conclusion that the ferment is elaborated by the glands, not as such, but as a zymogen, which is then transformed into a milk-curdling ferment by the action of hydrochloric acid. For the detection of this antecedent substance (zymogen), *Klemperer's* method may be employed:—To 2 cc. of filtered gastric juice are added 10 cc. of milk containing 2 cc. of a 3 per cent. solution of chloride of calcium, and an excess of a 1 per cent. solution of carbonate of soda. The mixture is then placed in an incubator, and if the zymogen be present, coagulation gradually ensues. The amount of the ferment formed varies chiefly with the quantity of hydrochloric acid generated in the process.

Experience has shown that the milk-curdling ferment is in excess in conditions of hyper-secretion and undue acidity of the stomach. When hydrochloric acid is wanting or but scantily present, the ferment is also absent or reduced in quantity; but according to *Klemperer*, the ferment may also result from its zymogen in presence of organic acids.

3. Acids.—The gastric juice contains hydrochloric acid, and also butyric, acetic, and lactic acids.²⁸

(a.) **ACIDITY.**—In very rare instances an increased quantity of acid has been found in the stomach,²⁹ and with this is sometimes coupled an excessive secretion of gastric juice.

*Riegel*³⁰ made the important discovery that, *in cases of round ulcer of the stomach, the acid constituent of the gastric juice is greatly in excess*; and the fact has been further established by the observations of *Korcynski, Jacorski*,³¹ and many others (see p. 175).

According to *Reichmann, Riegel, Sticker*,³² and others, a distinction is to be made between hyper-acidity and excessive secretion of the gastric juice. By attending to this point it should be possible better to discriminate between certain affections of the stomach, and especially amongst those conditions which are still classed under the general heading of *gastric catarrh, dyspepsia, and the like* (see p. 177).³³

A diminished acidity of the gastric juice occurs temporarily when a large quantity of alkaline substances has been swallowed, and as a persistent condition apparently in all febrile diseases.

The acidity of the gastric juice may be measured in this way:—A certain quantity—increased, if necessary, by the addition of water—is filtered, and its reaction tested. If this be acid, a known quantity of the filtrate is taken and coloured with a little neutral tincture of litmus. Solution of soda of definite strength (the 1-10 normal or deci-normal

soda solution may be used with advantage) is now added slowly from a graduated burette, until the point is reached finally at which the onion-red colour of the fluid gives place to a violet hue. From the quantity of soda used, that of the acid present may be known, 1 cc. of the normal soda solution employed in this way corresponding to 0.0365 grm. hydrochloric acid.³⁴ Instead of litmus an alcoholic solution of phenolphthalein may be used. A few drops of this should be added to the fluid before titration, and the alkali slowly supplied until a red colour begins to develop.

The conclusions arrived at by this method are correct only when the gastric juice contains hydrochloric acid alone, and not, as is generally the case, several other acids as well (see below). On this account *Ewald*³⁵ points out the fallacy of directly deducing the total acidity from the amount of a deci-normal solution of soda required to neutralise it. The expression 50 per cent. acidity implies that 50 cc. of the deci-normal solution of soda will neutralise 100 cc. of the gastric juice experimented upon.

To determine whether the acidity is due to the presence of free acid or to acid salts, resort may be had to the methods of *Uffelmann* and *Leo*, by which only *free* acid is detected. *Leo*³⁶ uses calcium carbonate, which in presence of acid is decomposed without heat, carbonic acid being given off and the fluid acquiring a neutral reaction. If no free acid, but only acid salts, be present, the fluid remains acid and reacts to litmus paper as before. To carry out the test a quantity of the gastric juice under examination is rubbed up with chemically pure calcium carbonate, and the reaction obtained before and after the addition of the salt is compared. If in the second case this be neutral, the original acidity was due to free acid; if, on the other hand, it be still acid, but less so than formerly, the fluid contained both acid salts and free acids. By an application of the same process it is possible to estimate the quantity of free acid—hydrochloric and organic acids—in a particular specimen. By *Leo*³⁷ this is done as follows:—10 cc. of the gastric juice is filtered, and 5 cc. of a concentrated solution of chloride of calcium, and a few drops of an alcoholic solution of phenolphthalein, are added. The mixture is titrated with deci-normal alkaline solution. Next, to 15 cc. of the filtered juice is added 1 grm. of dry powdered calcium carbonate, and the mixture is treated in the way described above. It is then passed through an ash-free filter—asbestos serves well—and the aspirator may be used to expedite filtration. Ten cc. of the filtrate are measured out and placed in a small flask. The stopper of this flask is perforated by two openings. Through one of these a glass tube passes to the bottom of the flask; the other transmits one end of a short right-angled glass tube, which reaches only just within

the flask, while the other end tapers a little and is connected by a caoutchouc binder with a Böhm's air-pump. By means of the latter the carbonic acid which forms is drawn off. The fluid in the flask is then treated with 5 cc. of calcium chloride solution and a few drops of phenolphthalein, and titrated. The difference between the results of this and of the former titration expresses the amount of acidity which is due to the presence of free acid. Should it be ascertained, in the way which will be described presently, that there were no organic acids in the fluid, the difference depends upon the hydrochloric acid alone, and its amount may be determined by the formula already given, namely, that 1 cc. of the deci-normal solution of soda corresponds to 0.00365 grm. of hydrochloric acid. This method is sound in principle and will be justified by its results. It must, however, be mentioned that exception is taken to it by *A. Hoffmann* and *A. Wagner*³⁸ on theoretical grounds. According to *Kossler*,³⁹ the process of filtration may be omitted, and the conclusions both as to acidity and to the proportion of free acid (see below) will be accurate.

(b.) **HYDROCHLORIC ACID.**—The gastric juice secreted during the later stages of digestion appears normally to contain only free hydrochloric acid. At an earlier period lactic acid is also present.

A. Detection of Free Hydrochloric Acid.—The examination of the gastric juice for free hydrochloric acid is attended with much difficulty, since the chlorine salts yield nearly all the same reactions as the free acid. To obviate this, many expedients⁴⁰ have been suggested; but we shall notice here only those methods which will serve for clinical purposes.

1. **Mohr's Tests.**⁴¹—(a.) To the gastric juice to be tested is added first a solution of iodide of potassium and starch-paste, and then a few drops of a very dilute solution of ferric acetate. If free hydrochloric acid be present, a blue coloration (starch iodide) appears. This very simple test is not altogether to be relied upon, inasmuch as it will yield a negative result in presence of phosphoric acid and its salts, even though free hydrochloric acid be present also.

(b.) The following test, also discovered by *Mohr*, answers its purpose admirably:—It depends upon the fact that a very dilute solution of ferric acetate, free from alkaline acetates, is unchanged by the addition of a few drops of sulphocyanide of potassium solution, and retains its yellow hue, while, if a mineral acid be present, it colours a deep red.

*Ewald*⁴² has obtained good results by applying the test in the following manner:—Two cc. of a 10 per cent. solution of sulphocyanide of potassium and 0.5 cc. of a neutral solution of ferric acetate are made up to 10 cc. (with water). A few drops of the solution, which is of a ruby-red colour, are placed in a small porcelain dish, and one or two drops of the fluid to be tested are allowed to trickle slowly on to it. If hydrochloric acid be present, a light violet colour forms at the point of contact of the two fluids, which gives place to a deep mahogany-brown when they mix. This test, according to *Ewald*, has an advantage over the aniline-dye tests in that its result is not materially affected by salts or peptone; but it is certainly less sensitive than the methyl-aniline-violet and tropæolin tests.

2. The Aniline Dye-Tests. (a.) **Methyl Aniline-Violet Reaction.**—

This reagent was first used by *Witz* and *Hilger*⁴³ for the detection of free mineral and organic acids. *Maly* has employed it for physiological, *Van der Velde* for clinical purposes.⁴⁴ To obtain the reaction, the fluid to be tested is mixed with a violet-coloured watery solution of methyl-aniline-violet. If very much free hydrochloric acid be present (as is never the case in the gastric juice), the fluid will be bleached. If a moderate quantity, it becomes green; and if very little, of a blue colour. The direct examination of the gastric juice never shows more than the transition from violet to blue. For the detection of a very small proportion of acid, *Maly* recommends that the mixture should be evaporated to the bulk of one or two drops on the water-bath. So little as $\frac{1}{2}$ mgrm. of hydrochloric acid will then cause the change from violet to blue.

*Kost*⁴⁵ recommends the addition of a 10 per cent. solution of tannin before testing with methyl-violet, in order to precipitate peptones, which would otherwise hinder the reaction.

(b.) **Tropæolin (oo)** in alcoholic or watery solution yields a ruby-red or dark-brown red colour in presence of free acids. *Ewald*⁴⁶ maintains that this reaction constitutes the most sensitive test for free lactic as well as hydrochloric acid. *Boas*,⁴⁷ who takes the same view, employs a tropæolin test-paper for the purpose.

(c.) **Fuchsin.**—The test with fuchsin is far from sensitive, and on that account of little utility.

(d.) **Emerald-Green.***—The so-called "crystallised" emerald-green affords a sensitive test for free hydrochloric acid. Concentrated solutions of hydrochloric acid give reddish brown, and very dilute solutions a grass- or a yellowish green colour, with this reagent.

A brilliant-green, obtained from the same laboratory, has proved a very efficient test, as shown by observations made by *Hellstrom* in the author's clinic. Five mgrns. of this reagent will serve to detect 0.48 mgrm. of hydrochloric acid dissolved in 6 cc. of water, giving to the solution a bright green tint. It is to be noted, however, that a similar effect is obtained where acetic, formic, or lactic acid is present in a higher degree of concentration. *Bourget*⁴⁸ also uses a brilliant-green.

* The other emerald-greens produced by *Bayer*, and distinguished as emerald-green (extra crystallised) and emerald-green ii. and iii., proved useful, but less sensitive. Of the other reagents tested, Kaiser blue (*Gärtner*, Berlin) was but little sensitive. Its solutions turned a brown-green with concentrated hydrochloric acid, and an azure-blue with dilute acid. A number of green pigments, prepared by *Poirier* of Paris, were ineffective as tests for the acid.

* This substance is made at *B. Bayer*'s laboratory, Elberfeld, and, with other reagents, has been made the subject of experiment at the author's request by *Dr. Voigt*, on whose authority the statement in the text is given.

*Köster*⁴⁹ has recently employed malachite-green with good results as a test for hydrochloric acid.

(e.) **Congo-Red.**—This is an aniline dye, which was first used as a test for free acid by *Herzberg*. It may be employed most conveniently in the form of filter-paper saturated with the reagent, as recommended by *Hösselin*, *Riegel*, and his pupils,⁵⁰ for the detection of free hydrochloric acid.⁵¹ This, when immersed in a fluid containing free hydrochloric acid, turns blackish-blue or blue, according as much or little acid is present. This effect is not obtained with organic acids or acid salts in dilute solutions, and the intensity of the reaction is lessened in presence of proteids and of salts in large proportion. The efficiency of the test is undoubted, and, notwithstanding that its use is subject to certain fallacies,⁵² the Congo-red test-papers must be classed with benzo-purpurin and the aniline-violet reagent as most suited to the purposes of the practitioner.

(f.) **Phloro-Glucin and Vanillin.**—The reagent recommended by *Günzburg*⁵³ contains 2 grms. of phloro-glucin and 1 grm. of vanillin dissolved in 100 parts of alcohol. When hydrochloric acid is added to this, it deposits beautiful red crystals. For the detection of the acid in the gastric juice it is employed thus:—To the fluid to be tested for acid an equal quantity of the reagent is added, and the mixture evaporated on the water-bath. The presence of hydrochloric acid is shown by a delicate rose-red tinge on the surface of the porcelain dish. In this way so little as 0.06 per cent. of the acid is discernible, and the reaction is not impeded by organic acids, albumin, or peptone. By its means the author⁵⁴ has often detected 0.001 mgrm. of acid in 10 cc. of gastric juice. It is further commended by *Haas*.⁵⁵ This observer has similarly employed other colour substances, as eosin and methyl-orange, but experience does not justify their use.⁵⁶ *Boas* and *Puriz*⁵⁷ have recommended resorcin for the purpose, but it is less sensitive than *Günzburg's* reagent.

(g.) **Benzo-Purpurin.**—A very sensitive colour-test is that furnished by benzo-purpurin 6 B. Five mgrms. will serve to show 0.39 mgrm. of acid dissolved in 6 cc. of water (*Hellström*), causing the dark-red colour of the solution to give place to a light violet. A similar change is effected with acetic, formic, and lactic acids; but the colour obtained with organic acids is rather a brownish-violet, and requires a greater quantity of the latter for its production; in the case of acetic acid, not less than 0.84 mgrm. Test-papers may be prepared by soaking strips of filter-paper in a saturated watery solution of benzo-purpurin 6 B, and subsequently allowing them to dry. If one of these be placed in the gastric juice, it will immediately stain a dark blue, provided hydrochloric acid be present in a proportion not less than 0.4 grm. to 100 cc.

A brownish-black tint may be due to the presence of organic (lactic or butyric) acids, or to admixture of these with the hydrochloric acid. The ambiguity in this case may be dispelled by placing the paper so stained in a test-tube and shaking it up with sulphuric æther, when so much of the colour as is due to the presence of organic acids will speedily disappear, leaving a lighter stain, or restoring the paper to its original tint. If hydrochloric acid alone be present, no change will be effected in this way, and even after the lapse of twenty four hours the blue stain will be only slightly displaced. It is important, of course, that the æther used should itself be free from acid. To ascertain this its reaction may be tested with blue litmus paper.

The action of the benzo-purpurin test is not seriously interfered with by peptone and serum-albumin, even when these bodies are present in large quantity, and acid salts have no effect upon it.

The following experiments are of interest, as showing the various effects obtained from benzo purpurin with hydrochloric and organic acids —

Two solutions were made, one of 4 grms. hydrochloric acid in 100 cc. of water, and another of 0.1 grm. benzo-purpurin 6 B in 600 cc. of water. On mixing together 3 cc. of each, a beautiful blue colour inclining to violet developed, and a coloured flocculent precipitate formed on standing. The addition of hydrochloric acid caused this precipitate to dissolve, and it re-formed on the further addition of the dye.

The same effect was produced whether the solution contained 0.4 or 0.04 grm. hydrochloric acid in 100 cc., 3 cc. being taken in each case. Three cc. of a solution holding 0.004 grm. hydrochloric acid, when added to 3 cc. of a solution of 0.1 grm. benzo purpurin in 600 cc. of water, gave an evident violet coloration with slight turbidity.

With formic or butyric acid, to obtain the reaction rather less than 0.04 grm. in 100 cc. of water was required; with acetic acid, something more than 0.04 grm.; with lactic acid, over 0.004 grm. in 100 cc. In all cases alike 3 cc. were taken of each solution.

A comparison of the Congo-red and benzo-purpurin 6 B test-papers shows that the latter are the more sensitive, and they deserve the preference for practical purposes. Hyper-acidity and the preponderance of organic acids in the gastric juice can be shown by this simple procedure in the space of a few minutes.

None of these coloration processes give entirely satisfactory conclusions. In cases where the reaction is positively obtained, free hydrochloric acid is undoubtedly present; but we may fail to obtain the result when the gastric juice contains albumin, peptone, or salts in considerable quantity, even when free hydrochloric acid is present also.⁵⁴ The reactions with methyl-aniline-violet, Congo-red, phloro glucin and vanillin, and benzo-purpurin are the most to be depended upon. They will not serve for scientific purposes, but in view of their simplicity they are of the utmost value in bedside observation.⁵⁵

3. Uffelmann's Tests.—*Uffelmann*⁶⁰ has employed the colouring matter of claret in testing for free acids in the gastric contents, and quite recently, as a still more sensitive reagent, the amylic alcohol extract of bilberries, which he applies by means of blotting-paper soaked in it.⁶¹ The reaction depends upon the fact that the colour of such a test-paper changes in presence of hydrochloric acid even when peptone, albuminates, and salts are present, from greyish-blue to a rose tint, which persists after the paper has been washed with æther.

Lactic, acetic, and butyric acids give similar reactions, but only when in such a degree of concentration as is never found in the gastric juice; and, moreover, the reaction obtained with them is destroyed by the addition of æther.

[*Dreschfeld*⁶² employs Uffelmann's test in a modified form. The test solution consists of 0.5 cc. of claret (unadulterated), 3 cc. of 90 per cent. alcohol, and 3 cc. of æther; the solution is almost colourless, and is rendered a rose colour by the presence of a minute quantity of hydrochloric acid. This test is said by *Dreschfeld* not to be interfered with by the presence of peptone or albumin. Lactic acid gives a similar reaction only when occurring in a more concentrated form. The mixture does not keep long, and has to be freshly prepared.]

4. Ultramarine and Zinc Sulphide.—These substances were suggested by *Maly*, and employed by *Kahler*⁶³ as a test for free hydrochloric acid in the contents of the stomach. Ultramarine, according to *Kraus*,⁶⁴ is a test for free acids in general. It is decomposed by them even in dilute solutions, sulphuretted hydrogen being given off, while silicic acid and sulphur are precipitated. Zinc sulphide, again, is dissolved in dilute acids with the evolution of sulphuretted hydrogen. It is, however, insoluble in acetic acid.

In testing for hydrochloric acid the process is as follows:—About 20 cc. of the fluid under examination is placed in a crystallising crucible, and so much ultramarine is added as will suffice to give it immediately a blue tinge. The crucible is then covered with a watch-glass, from which depends a strip of filter-paper soaked in solution of sugar of lead, and the mixture is gently heated in the water-bath. After the lapse of a quarter of an hour, if hydrochloric acid be present, the blue colour of the fluid will have given place to a brown tint, while the lead-paper will be stained brown or black. Sulphide of zinc (as much as will fit on the point of a knife) is then added to another specimen, and the same process repeated, when the brown or black stain upon the lead-paper will again show the presence of hydrochloric acid. The reactions are rendered more feeble by the presence of salts and of phosphates in particular. They can be obtained also with organic acids (lactic and acetic) in more concentrated solutions.

These circumstances, and the comparative complexity of the process, render its application at the bedside a matter of difficulty. Whereas, on the other hand, we possess in the methyl-aniline-violet, phloro-glucin and vanillin, Congo-red, benzo-purpurin, and brilliant-green reactions a series of tests which are at once ready and accurate.

B. Quantitative Estimation of Free Hydrochloric Acid.

This can be accurately effected by the very complicated process of *Bidder* and *Schmidt*.⁶⁵ All the acids and bases in the gastric juice are quantitatively estimated, the proportion of each in 100 cc. of fluid ascertained, and their equivalents computed. The remaining hydrochloric acid is that which is free in the secretion.

Another method for the determination of this body depends upon the fact that the acid is insoluble in æther, whilst organic acids are soluble in that medium. To utilise this property for the purpose in hand, *Richel*,⁶⁶ following *Berthelot*, shakes up the gastric juice with æther, and determines by titration the

quantity of acid which is taken up by the latter together with what is retained in the watery solution. *V. Moracewski's* method⁶⁷ is based on the same principle.

Recently, *v. Mering* and *Cahn*⁶⁸ have adopted the expedient of collecting the volatile acids by distillation, lactic acid by extraction with æther, and combining the hydrochloric acid separated from the organic acids with cinchonine, shaking up the newly-formed hydrochlorate of cinchonine with chloroform, changing the acid into its silver salt, and finally weighing the chloride of silver obtained. *Köster* (see p. 157) has endeavoured to determine the quantity of hydrochloric acid in the gastric juice by a process of titration with alkalies after the addition of methyl-aniline-violet.

Günzburg's reagent, according to *Ewald*,⁶⁹ will also furnish a means of approximately estimating the quantity of hydrochloric acid present.

1. Leo's Method.—This has been already described (*vide supra*). If fatty acids and lactic acid be present, their proportion must be determined (*vide supra*), and deducted from the total acidity. The difference will express the quantity of HCl. According to *Kossler*,⁷⁰ the method is accurate. Both it and that which follows serve well enough for the estimation of physiologically active hydrochloric acid.

2. Sjöqvist's Method.—*Sjöqvist*⁷¹ has recently introduced a process for the estimation of free hydrochloric acid in the gastric juice founded upon the following facts:—The acids of the secretion may be changed into their barium salts by the action of barium carbonate, and when these are incinerated, the baryta salts of the organic acids leave barium carbonate, whilst the chloride of barium resulting from the combination with hydrochloric acid remains unchanged. The latter may then be separated from the insoluble carbonate by extracting the ash with warm water, and its quantity estimated by titration with chromate solution. The details of *Sjöqvist's* method are these:—Ten cc. of the gastric juice are filtered and placed in a platinum or silver crucible, and barium carbonate *free from chlorides* added in excess. The fluid is then evaporated to dryness at a gentle heat, and the residue charred and strongly heated for some minutes. After cooling, the residue is treated with 10 cc. of water, the mixture rubbed up, extracted repeatedly with boiling water, and filtered until the filtrate has a bulk of 50 cc. The quantity of chloride of barium in solution is best estimated by titration with bichromate of potash. This body gives with salts of barium a precipitate of barium chromate, which is insoluble in water and acetic acid, and soluble in hydrochloric acid. A solution of bichromate of potash of known strength is added from a burette, until all the barium present is precipitated in the form of chromate. A subsequent excess of bichromate of potash would give to the fluid a deep red colour, which would tend to mask the result. This may be prevented by the use of tetra-paper (tetramethylparaphenyl-diamine), which has the property of staining blue with oxidising substances. In the process of titration, therefore, the filtrate is mixed with one-

fourth or one-third its volume of alcohol and 3–4 cc. of a solution holding 10 per cent. acetic acid and 10 per cent. acetate of soda, and titrated with a solution of bichromate of potash (8.5 grms. to the litre) until a faint trace of blue appears upon the test-paper. The addition of acetic acid and acetate of soda has for its object to promote the precipitation of chromate of barium, and at the same time to prevent the formation of chromate of lime from the small quantity of lime salts and free hydrochloric acid that may be present. From the quantity of bichromate of potash used, that of the barium salt formed, and also of sulphuric acid present, result directly.⁷²

This process is attended with difficulty, and it is open to the objection that it depends too much on the judgment of the observer. The following modification of it is more accurate.

3. R. v. Jaksch' Modification of Sjöqvist's Method.—It is the author's practice to convert the chloride into barium sulphate, and by weighing the sulphate to calculate the amount of hydrochloric acid in 10 cc. of gastric juice. To that end the unfiltered gastric juice (10 cc.) is treated with chlorine-free carbonate of barium in excess, and placed on a furnace in a thin porcelain crucible, where it is evaporated to dryness, then gently fused in a muffle; the residue cooled, extracted with boiling water, and filtered; the filtrate evaporated on the water-bath to a volume of 100 cc., and dilute sulphuric acid added. The precipitate (sulphate of barium) is placed on a thick ash-free filter, washed with water, fused in a platinum capsule, thence removed with the usual precautions.⁷³ The result is calculated thus:—233 parts by weight of barium sulphate (BaSO_4) correspond to 73 parts of hydrochloric acid (HCl). And the quantity of the latter contained in 10 cc. of the gastric juice may be calculated from the formula

$$x = \frac{73}{233} \times M = 0.3132 \times M$$

where M = the quantity of barium sulphate obtained from 10 cc. gastric juice

x = the quantity of hydrochloric acid sought in 10 cc.

This method enables the examination to be effected within a comparatively short time. Its accuracy is attested by *Leo* and *Leubuscher*.⁷⁴ The objections made to it on the ground of its being too complicated are ill-founded.⁷⁵ It serves for the estimation of HCl equally when free and when combined with organic digestive products (proteids). It is doubtless true that there are proteids which enter into combination with HCl in such a way that the acid is no longer perceptible by this process,⁷⁶ but the author's investigations have satisfied him that such combinations do not in fact occur in digestion. Quite lately *Leo*⁷⁷ has contended against the principle of *Sjöqvist's* method, adducing considerations which

gravely affect the pretensions to accuracy both of that method and of the modification of it just described ; the result of *Kossler's*⁷⁶ researches has added to the force of *Leu's* objections. It would appear, indeed, that the method is inapplicable where phosphates are present, and its utility is much curtailed by this fact. In any case, it is only the absolute values as determined by the method which are affected. The conclusions based upon them remain good. Finally, the observations of *Rosenheim*⁷⁹ show that the fallacies which *Leu* has pointed out do not apply to a bedside examination, and *v. Pfungen*⁸⁰ maintains the utility of the method for clinical purposes. The modifications of *Salkowsky* and *Fawcitzky*⁵¹ and of *Boas*⁸² offer no special advantages. Similarly, *Mierzinsky's*⁸³ proposal to estimate the hydrochloric acid by volumetric analysis is, as *Wiener*⁸⁴ has shown, impracticable. Of *Bourget's* process⁸⁵ the author has no experience. That of *Winter* and *Wagner*, according to *Kossler*,⁸⁶ yields an estimate of HCl, free and combined with proteids, which is somewhat too high. *E. Biernacki* and *L. Sansoni*⁵⁷ assert that the results obtained by the *Hayem-Winter* method are inaccurate.

4. Braun's Method.⁸⁸—A certain quantity—5 cc.—of the filtered gastric juice is taken, and its acidity determined by titration with $\frac{1}{10}$ normal soda solution in the manner described at p. 153. To another 5 cc. of gastric juice is added soda solution a little in excess of what was needed to neutralise it. The fluid is now incinerated (see p. 160), and to the ash is added as many cc. of $\frac{1}{10}$ normal sulphuric acid solution as were needed of $\frac{1}{10}$ normal soda solution to neutralise the specimen taken, i.e., 5 cc. of the filtered juice. The ash is thus dissolved; the fluid is warmed, and carbonic acid driven off, after which a solution of phenolphthalein is added to it, and it is titrated with $\frac{1}{10}$ normal alkali solution. The number of cc. of $\frac{1}{10}$ normal soda solution employed, multiplied by 0.00365 (see p. 154), gives the quantity of HCl in 5 cc. of gastric juice. This method is founded on the same principle as *Sjögren's*, but, according to *Kossler*,⁸⁹ it is not accurate, since the acidity which is due to acid phosphates is not allowed for.

5. Hoffmann's Method.⁹⁰ In testing the proportion of HCl in the gastric juice, *Hoffmann* has availed himself of the property which HCl possesses of inverting cane-sugar, i.e., of breaking it up into dextrose and levulose, so that the polarisation-phenomena of its solutions are altered. The following preparations are required. 1. A fluid containing known quantities of cane sugar and HCl. 2. Equal quantities of cane-sugar and gastric juice. 3. Gastric juice alone. 4. Gastric juice with cane-sugar and sodium acetate in equal quantities. The rotatory power of each of the four fluids is ascertained by means of the polarimeter, and they are then allowed to stand in a warm place for some hours, and their rotatory power again investigated. The calculation is then made by the formula, $\log A - \log (A - x) = C$, where A = the quantity of sugar originally present, x = the quantity which has been converted at the termination of the process. This method is undoubtedly ingenious, but it is subject to the drawback that it

requires a very accurate polarimeter, eight polarimetric examinations, and a highly-complicated calculation. Recently it has been much simplified by substituting titration with methyl acetate for inspection with the polarimeter.⁹¹ The researches of *Kossler*⁹² have shown, however, that it serves only for the estimation of free HCl, to the exclusion of that which is combined with proteids.

In addition to those described here, many other methods have been brought forward for the estimation of HCl, but they possess no superior advantages, such of them as are easier of application being proportionately wanting in accuracy. Amongst them are those of *C. Th. Mörner*, *Mintz*, *Jolles*, *Kronfeld*, *Czernianski*, and *Töpfer*.⁹³

6. Lüttke's Method.⁹⁴—Ten to 20 cc. of gastric juice are taken, and the estimation of total chlorides is made as described in Chapter VII. Next, 10 cc. of gastric juice are dried and gently fused. A comparison of the results of these two processes gives the quantity of hydrochloric acid.

The researches of *Martius* and *Lüttke*⁹⁵ are of importance as showing that lactic acid is not a normal product of digestion (see p. 155). Further experience is needed before an opinion can be expressed as to the utility of this method for the purpose of estimating hydrochloric acid.

(c.) THE QUANTITY OF HYDROCHLORIC ACID PHYSIOLOGICALLY ACTIVE IN THE GASTRIC JUICE, AND ITS DIAGNOSTIC IMPORT.—Concerning the quantity of hydrochloric acid which is secreted normally during digestion the recorded observations are very few. *Moritz*, *Wohlmann*, and *v. Jakob*⁹⁶ have investigated this subject. According to the latter, the quantity formed during digestion in healthy children varies greatly with the nature of the food, and generally attains its maximum within one to three hours after a meal. With milk, which combines very readily with acids, the increase is slow; it is more rapid with nitrogenous, slowest, but with greatest initial rapidity, with farinaceous food. The greatest quantity of effective HCl was obtained with a diet of milk alone, a smaller quantity with a meat diet, and the least with carbohydrates. The quantities were respectively: 0.1615 grm. (mean of fourteen observations), 0.1563 grm. (mean of eleven observations), and 0.1102 grm. (mean of ten observations), in 100 cc. of the gastric contents. The facts are the same in healthy adults. Thus with the method described at p. 161, the author has found that when 200 grms. of ham have been taken, there are in 100 cc. of the gastric contents 0.0643 grm. of HCl in thirty minutes, 0.1529 grm. in forty-five minutes, and 0.0992 grm. in an hour. From this it follows that, as a preliminary to basing any inference upon the quantity of HCl secreted, it is necessary to consider what food the subject of the inquiry has taken, and at what time he has taken it. *The absence of free HCl, or its presence only in very small quantity, fifteen*

to thirty minutes after a meal, has no pathological significance. But should there be little or no free HCl present one to three hours after taking milk or nitrogenous food, the fact is evidence of a grave defect of function. A large quantity of HCl, even so much as 0.33 per cent. three hours after food, does not necessarily imply functional disorder (hypersecretion). Such considerations must always be weighed in forming an inference for diagnostic purposes. Again, for practical purposes, those tests alone are satisfactory which yield information concerning the physiologically-effective acid. From this point of view the colour-tests are insufficient, but they have the advantage of being easily applied, and where approximate results are desired they serve well enough. For scientific purposes the requirements are:—1. The application of such methods as dispense with the necessity of filtering the gastric juice, since this process is attended with much waste of the acid (*r. Jaksch*).⁹⁷ 2. That the method chosen should be one which takes account of that part of the acid which is physiologically effective. These requirements, as *Kossler* has shown, are fulfilled only by *Leo's* method, when applied to artificial digestion. Whether this is true of the natural gastric juice also remains to be proved. Approximately accurate results, however, may be obtained by the author's modification of *Sjögqvist's* process. One of these methods may be used with advantage to control the other. We shall return to this point presently in treating of the contents of the stomach in different gastric disorders.

It may be suggested here that for the terms "free" and "combined" hydrochloric acid, "physiologically active" and "physiologically inactive" should be substituted. By the first would then be meant either that portion of the acid which has already discharged its function and has entered into combination with proteids, or that which is still available, and therefore in the literal sense free.⁹⁸

The investigation of the functions of the stomach in disease of all kinds, and especially with reference to the secretion of hydrochloric acid, has of late years been pursued with the utmost energy. The contributions to the subject which possess the chief diagnostic interest may be briefly mentioned. *Immermann* and *Schetty*⁹⁹ found that in tuberculosis there was no change in the secretion of HCl. Their conclusions are supported by *Chelmonski*, *Klemperer*, *O. Brieger*, *Hildebrand*, and *Schwalbe*.¹⁰⁰ *Grusdew*,¹⁰¹ on the other hand, observed a diminished production of the acid. *Hüffler*¹⁰² states that in heart-disease the acid is deficient, but this is not in accordance with the observations of *Einhorn*, *Adler*, and *Stern*.¹⁰³

*Biernacki*¹⁰⁴ and the author have noticed a considerable deficiency of the acid in renal disease in many cases.¹⁰⁵ *Lenhartz*¹⁰⁶ has collected much information upon this subject. In acute and chronic dyspepsia

there was a remarkable deficiency of free acid; in chlorosis a similar deficiency was observed in 45.6 per cent. of the cases investigated, whereas in gastric ulcer the condition was inconstant. *Geigel* and *Abend*¹⁰⁷ obtained very varying results in cases of neurotic dyspepsia.

From these facts it results that the presence or absence of free hydrochloric acid is a symptom of doubtful import, and that it must be weighed in conjunction with the other circumstances of the case. It is much to be desired that measures should be taken for the acquirement of accurate data concerning the production of hydrochloric acid in diseases of the stomach and other parts, and this may be done by the use of the more scientific methods indicated here, and especially by observance of the precautions mentioned on p. 163. It is sufficient here to point out that a failure of the secretion on the one hand, and its production in excess on the other, are alike evidence of disease.¹⁰⁸ Their precise significance will be dealt with later (pp. 174, 178).

(d.) **ORGANIC ACIDS OF THE GASTRIC JUICE.**—In this connection we have to deal with lactic, acetic, and butyric acids.

1. **Lactic Acid.**—*A. Qualitative Tests.*—For the detection of this body in the gastric juice the carbolo-chloride of iron test is to be recommended (*Uffelmann*¹⁰⁹), (*Kredel*).¹¹⁰ To a mixture of 10 cc. of a 4 per cent. solution of carbolic acid with 20 cc. of water, a few drops of perchloride of iron solution are added, and the resulting amethyst-blue colour changes to yellow in presence of a few drops of lactic acid. Alcohol, sugar, and phosphates yield a similar reaction (*Ewald*);¹¹¹ but where the colour-change is quickly and markedly effected, it shows the presence of lactic acid, and in this case furnishes a useful clinical test.

*Boas*¹¹² recommends the following process:—Oatmeal broth is taken, and the stomach contents removed and filtered; 10–20 cc. of the filtrate is evaporated on the water-bath to a syrupy consistence, after the addition of carbonate of baryta in excess should the tests by Congo-paper, &c., show the presence of free acid. To the thick fluid are added a few drops of phosphoric acid; the carbonic acid formed is driven off by heat, the fluid again allowed to cool and extracted with 100 cc. of alcohol-free æther. After half-an-hour the clear æther is poured off, evaporated, and the residue dissolved in 45 cc. of water, which is well shaken up and filtered. The filtrate is treated with 5 cc. of sulphuric acid (sp. gr. 1.84) and a little manganese. The fluid is then placed in an *Erlenmayer's* flask with a stopper through which passes a bent tube of glass, the longer limb of which is led into a glass cylinder containing 5–10 cc. of alkaline iodine solution or *Nessler's* reagent, and the flask is heated. The lactic acid present yields aldehyde, which combines with alkaline iodine solution to make iodoform, or with *Nessler's* reagent to produce a reddish-yellow mercury-aldehyde. The author would notice

that Boas, in this contribution, claims to have found that aldehyde responds to Reynolds' acetone test; but would point out that he discovered this peculiarity in aldehyde nine years earlier.¹¹³

A further test for this body is derived from a very dilute solution of perchloride of iron—two to five drops of a watery solution of perchloride in 50 cc. of water.¹¹⁴ The faint yellow colour of the fluid, whilst not affected by the addition of hydrochloric, butyric, or acetic acid, is intensified in presence of dilute lactic acid. To separate lactic acid from the gastric juice, the distillation residue (see below) of the gastric juice, in which the acid is dissolved, may be extracted with æther, and submitted to the tests described elsewhere (see chapter on *Urine*).

Many observers apply the tests for lactic acid not directly to the gastric juice, but to the residue of an æthereal extract made from it.

B. Quantitative Estimation.—This may be effected by *Cahn* and *v. Mering's* method (see p. 160), or by that of *Leo*.¹¹⁵ Ten cc. of gastric juice are taken, and, when the fatty acids have been removed (see below), extracted six times with 100 cc. of æther in a separator-funnel, the resulting æthereal extracts collected, the æther driven off by exposure to the air by heat from a water-bath—(a flame must not be used)—and the residue dissolved in water. The acidity of the solution is then determined by a $\frac{1}{10}$ normal soda solution. Since 1 cc. of the soda solution corresponds to 0.009 grm. of lactic acid, the quantity of the latter contained in 100 cc. of gastric juice may be obtained by multiplying the number of cc. of alkali used by 0.009.

Boas' test is the basis of an estimating process. The flask (see p. 165) is then furnished with a stopper which transmits a second tube of large calibre reaching within to the bottom of the fluid, and outside terminating in tubing secured by a clamp, and connected with a collapsing india-rubber ball. The fluid (p. 165) is now heated. Aldehyde passes over, and the last traces are expelled by a current of air forced through the second glass tube. The distillate is led into a well-stoppered Erlenmayer's flask containing 20 cc. of a $\frac{1}{10}$ normal iodine solution and 20 cc. of caustic potash (56 cc. potassium hydrate in 1000 cc. of water). The flask should be closed with the receiver, with some of the iodine solution in it. At the end of the distillation all the iodine solution is collected in the flask, stoppered and well shaken, and allowed to stand for a few minutes. Then there are added 20 cc. of dilute hydrochloric acid (sp. gr. 1.018) and an excess of bicarbonate of sodium; and finally so much of a $\frac{1}{10}$ normal sodium arsenite solution continuously supplied as will suffice to decolorise the fluid, and then, by titration, the point at which a permanent blue colour is obtained with a fresh solution of starch is sought. The number of cc. of $\frac{1}{10}$ normal iodine solution, less the quantity of arsenious acid used, gives

the quantity of iodine in combination as iodoform. One cc. of $\frac{1}{10}$ normal iodine solution represents 0.003388 grm. of lactic acid.

2. Butyric and Acetic Acids.—(a.) *Qualitative Tests.*—If the gastric contents be extracted with æther, butyric and acetic acids may be recognised by their smell (*Uffelmann*). To separate these acids, the gastric juice is distilled, and the distillate tested in the manner laid down for the examination of the urine.

*Hammarsten*¹¹⁶ prefers not to distil the gastric juice directly, but to neutralise it first with caustic soda, and then to extract with alcohol, proceeding afterwards in the manner to be described for the detection of fatty acids in the urine. The object is to avoid the error of including fatty acids derived from proteids.

*Uffelmann*¹¹⁷ directs attention to the importance of a systematic analysis of the gastric juice for the detection of free acids. To do this the contents of the stomach are filtered and their reaction tested. Should this be acid, they are submitted to the following process:—The total acidity is determined by titration with a deci-normal solution of caustic soda, and a portion is tested with dilute solution of perchloride of iron for the presence of lactic acid. Another portion is then tested for free hydrochloric acid with bilberry-dye test-papers. A rose colour obtained when the degree of acidity is slight, and persisting after the addition of æther, indicates the presence of hydrochloric acid. If, on the other hand, the colour is entirely destroyed by treatment with æther, it is evidence of considerable quantities of lactic, butyric, and acetic acids.

*Riegel*¹¹⁸ and *Koster* have employed similar methods with success; and attention may also be directed to the process described at p. 157.

(b.) *Quantitative Estimation.*—*Leo's*¹¹⁹ method is the following:—Ten cc. of the gastric juice are taken, and the total acidity determined in the manner described at p. 163. Another 10 cc. are filtered and boiled until the fumes no longer present an acid reaction. The residue is allowed to cool, and is then titrated with deci-normal solution of soda. The difference between the acidity of this and the former specimen is that due to the fatty acids. The method is not absolutely accurate, since HCl may be driven off by boiling.

4. Proteids.—Proteids occur in the gastric contents during digestion, being partly formed in that process, and in part derived from the food. Their recognition affords valuable evidence as to the functional condition of the stomach; and to make its import clearer, it will not be out of place to refer to certain facts in physiology. The period of digestion may be divided into two stages. (1.) The first of these, which lasts but a short time (15–20 min.), is occupied chiefly with the digestion of starchy matter, and is characterised by the presence of the resulting products, and especially lactic acid. (2.) The second stage commences

with the secretion of pepsin and an active gastric juice, by means of which the albumin of the food is changed. The two stages pass gradually into one another, and authorities are not agreed as to whether lactic acid occurs only during the first (*Ewald, Boas*, and others), or in the second stage of healthy digestion also, when it is said by some (*Cahn and v. Mering, Ritter and Hirsch*) to be present together with the more abundant hydrochloric acid. The observations of *Martius* and *Lüttke*, and of *Boas*,¹²⁰ seem to show that lactic acid does not occur in the stomach in appreciable quantity at any period of the digestion of food which does not contain carbohydrates. It would appear, moreover, that the one acid may replace the other in respect of its action upon proteids (*Feranini*¹²¹).

For the purpose of an examination, whether in a healthy individual or otherwise, a test-meal should be administered on an empty stomach. This, according to *Ewald*, should consist of a dry, well-baked roll, and water or weak tea; whilst *Leube* and *Riegel* recommend a meal of water-broth,* semolina and flour-gruel,† and meat. *Ewald's* regimen has the advantage that digestion is at its height within an hour after the food has been taken, whereas in the other case it is necessary to wait for four to six hours before the examination can be begun. The contents of the stomach are obtained in the manner described at p. 151. A test-meal of this kind is very useful for many purposes. *Klemperer* and the author¹²² administer milk in the same way. It is probably advisable that the test-meal should be of the simplest possible character, a single proteid, as, for example, egg-albumin, or a carbohydrate being given, and the choice will be made in accordance with the subject under investigation. The proteids in question are albumin, hemialbumose, peptone, and syntonin.

Albumin and *hemialbumose* may be detected by the process detailed in the chapter on *Urine*. Should these bodies and *syntonin* be absent, the biuret reaction (red coloration) will serve directly to show the presence of *peptone*. If, on the other hand, following the method referred to, other proteids (and especially those which are coagulable by heat) are found to be present, these must first be removed in the usual manner (see chapter on *Urine*), provided a sufficiency of material remains to work upon. The filtrate may then be submitted at once to the biuret test, the previous precipitation with phosphotungstic acid not being necessary.

Syntonin may be known by its being precipitated by neutralisation from its acid solutions.

* [*Wassersuppe*, translated here as water-broth, is made of boiling water with small squares of dry rolls, some salt and fresh butter.]

† [*Griessuppe*, semolina soup, consists of semolina boiled in water and seasoned with salt and butter or extract of meat.]

About 30-40 cc. of gastric juice will suffice for an examination of this kind when a little skill has been attained in conducting it. The pathological specimens which come to hand rarely exhibit other nitrogenous bodies than peptone, as is the case also when the contents of the stomach are examined several hours after the test-meal has been taken.

5. Carbohydrates.—Grape-sugar may sometimes be found in the stomach, having either been introduced with the food, or formed there by the action upon starch of saliva which has been swallowed. This latter mode of origin belongs especially to conditions of hypersecretion of HCl (*Riegel*, see p. 153, *Ewald*).¹²³ The mode of testing for sugar is the same here as in the blood (p. 87), the proteids being first removed.

The phenomena of the digestion of starch, and the formation of its intermediate products, involve some points of interest. An hour after food has been taken, under ordinary circumstances, neither starch (blue colour with iodine and iodide of potassium solution) nor erythrodextrin (red with the same reagent) can be discovered in the filtered gastric juice. Should it happen otherwise, some cause tending to delay the amylolytic process may be inferred, and this may be sought either in the fact that the saliva is deficient in diastase, or that there is an excessive secretion of free acid by the stomach at the outset of digestion (*Ewald*, *Boas*,¹²⁴ *Rosenheim*¹²⁵). In health, also, when amyloseous food has been taken in quantity, starchy particles may be found, and their nature determined chemically (see p. 172).

6. Urea.—For the detection of urea, one of the methods adopted for the same purpose in connection with the blood (p. 83) may be employed. Considerable quantities of this body are found in the stomach in cases of uræmia.

7. Ammonia.—Salts of ammonia are abundantly present in the stomach in rare instances. Where a considerable bulk of the gastric contents can be obtained (as by vomiting), the quantity of ammonia present may be estimated, after the removal of the proteids, by *Salkowski's* method.¹²⁶ For this purpose, 50 cc. of the vomited matter are taken, 20 grms. of pure powdered chloride of sodium first added, and then 100 cc. of a mixture holding seven parts by volume of a saturated solution of chloride of sodium and one part of acetic acid (1.040 sp. gr.). The whole is then mixed, and allowed to stand for fifteen to twenty minutes, when it is measured and filtered. Of the proteid-free filtrate 50 to 100 cc. are measured off, treated with milk of lime, and placed under a bell-glass containing a known quantity of 1/100 normal solution of an acid. After the lapse of from three to five days, the latter is removed and titrated with 1/100 normal alkali solution coloured

with rosolic acid. In this way the quantity of ammonia absorbed may be determined.¹²⁷ According to the later researches of *Rosenheim* and *Strauss*,¹²⁸ it would seem that the ammonia salts, which are for the most part derived from the food, are to some extent also a product of the gastric glands.

The same process is applicable to the determination of ammonia salts in the blood and other fluids. Still better for this purpose is the method of vacuum distillation employed by *v. Nencki* and others.

8. Potassium Cyanide.—*Kelling's*¹²⁹ observations seem to show that this salt occurs in the stomach. It may be tested for in the manner given in Chapter II.

9. Hydrogen Sulphide.—Recent researches have dealt with the presence of sulphuretted hydrogen in the gastric contents (*Bous*,¹³⁰ *Zawadski*¹³¹). It may be derived by communication from the large intestine in obstruction, or from the mouth in persons with dental caries. The eructations from gastric carcinoma are sometimes foetid with this gas. For its detection see Chapters VI. and VII.

10. Hydrogen and Carbonic Acid Gases.—On this subject *J. Hoppe-Seyler*¹³² may be consulted. The antecedents of these gases in the stomach he found to be very variable.

5. Estimation of the Rate of Absorption of the Gastric Contents.—The rapidity with which the gastric contents are absorbed, and, consequently, the functional activity of the stomach in this respect, may be determined thus, after *Penzoldt* and *Faber*.¹³³ A capsule containing 0.1 grm. of iodide of potassium is given to the patient to swallow. The saliva is then tested for iodine every two or three minutes, by placing a little of it upon filter-paper saturated with starch-paste, and adding a drop of fuming nitric acid. The presence of iodine is shown by a blue colour, which usually appears in 8–15 minutes. According to *Ziceifel*¹³⁴ this period is prolonged—thus indicating a deferred absorption—in various affections of the stomach, as dilatation, cancer, and gastric ulcer.

6. To Determine the Contractile Activity of the Stomach.—For this purpose various expedients have been adopted. Amongst them may be mentioned those of *Leube*, *Klemperer*, *Sievers*, and *Ewald*.¹³⁵ *Leube* assumes that the motor function of the gastric walls is impaired when the use of the sound shows the presence of food seven hours after it has been taken. *Klemperer's* method is objectionable on the score of discomfort. *Ewald* introduces within the stomach a gelatine-capsule containing 1 grm. of salol. This decomposes into phenol and salicylic acid as soon as it reaches the small intestine, and rapidly appears in the urine, where it may be detected by solution of ferric chloride (see chapter

on *Urine*). The interval which elapses before the drug appears in the urine may be taken to indicate the length of time required for the stomach to discharge its contents. In health this is from 40 to 60 minutes, but a much longer time in gastric atony and dilatation. The results obtained in this way have no utility in diagnosis.¹³⁶

7. A Summary of the Chemical Examination of the Gastric Contents.—It is seldom that sufficient material can be obtained for the systematic examination suggested here, and it will be necessary to make the investigation by successive evacuations of the stomach, the administration of test-meals, &c., before the different processes can be applied.

1. The reaction is to be tested.
2. A known quantity of the fluid, say 10 cc., is taken for the determination of acidity.
3. Another quantity of 10 cc. is examined to show the presence of pepsin and milk-curdling ferment.
4. The benzo-purpurin, Congo-red, and brilliant green tests for free hydrochloric acid are applied, and the latter estimated quantitatively by *Sjöqvist's* method.
5. A rough estimate is made of lactic, butyric, and acetic acid in the manner described at pp. 165-167.
6. Examination for proteids, this being confined to serum-albumin and peptone where sufficient material cannot be had.
7. Test for starch and its digestive products.
8. The remainder of the fluid is distilled, and the residue shaken up with æther, to determine accurately the quantity of lactic acid which it contains (p. 166). The distillate is tested for fatty acids in the manner described in the chapter on *Urine*.

II. THE INTESTINAL JUICE.—In the present state of our knowledge the investigation of the intestinal juice lends but little aid to clinical study. Notwithstanding this, the subject is worthy of attention, and it is probable that before long it will be brought within the scope of a practical inquiry.

1. Naked-Eye Characters.—The intestinal juice is a mixed secretion derived from several glands, and its character varies with the part of the tract from which it is taken. In the small intestine it is the product of Brunner's and Lieberkühn's glands, the liver, and the pancreas. It is only this fluid, which is a compound of bile, pancreatic fluid, and the secretion of the intestinal glands, that will be noticed here. It is a clear yellow thin fluid of a strongly alkaline reaction, and sp. gr. 1.009-1.011. On standing exposed to the air it turns to a grass-green (biliverdin) colour.¹³⁷

2. Formed Elements.—Concerning these nothing is accurately known.

3. To Obtain the Intestinal Juice.—According to *Boas*,¹³⁸ it is first ascertained whether the stomach is empty. Should it be so, the patient is made to lie down, and the abdomen over the region of the gall-bladder is massaged ; then the patient is made to stand upright, and the sound is again introduced, when he once more assumes the horizontal position, and the fluid is pressed out.

4. Chemical Constitution of the Intestinal Juice.—The intestinal juice contains bile acids and bile pigments, syntoinin, peptone, a small quantity of leucin and tyrosin (comp. Chapter VII.), and a number of ferments, of which the chief are the tryptic, fat-splitting, and emulsifying * (pancreatic), diastatic, and inverting ferments.

Concerning the changes which the secretion undergoes in disease nothing is yet known. *Boas* was the first to introduce the subject of digestion in the small intestine. Physiological research has yet to pave the way before our knowledge in the matter can be applied to the purposes of diagnosis. Nevertheless, the few facts which have been brought to light by *Boas* and *Noorden*¹³⁹ afford ample prospect of a rich harvest both of physiological and clinical results from further study in this direction.¹⁴⁰

III. EXAMINATION OF THE VOMIT.—The vomit includes the secretions of the mouth and nasal passages which have been swallowed, and are, for the most part, already undergoing digestion, the gastric juice and ingested substances, in part altered by the action of the stomach, and partly unchanged. Further, it sometimes contains bile.

The naked-eye and microscopical appearances vary with its constitution, and chiefly with the abundance and character of the food. Apart from such constituents as are derived from the mucous membrane of the mouth and nasal passages, and which have been already described, the vomit almost invariably presents—(1) Columnar and squamous epithelium, both usually much altered in form ; (2) isolated white blood-corpuscles, generally so transformed by the action of the gastric juice that little more than the nuclei remains ; (3) isolated red blood-corpuscles, usually seen as colourless rings, and very seldom in a quite perfect state ; (4) the following derived from the food :—

1. Muscle fibres, readily recognisable by their transverse striation.
2. Fatty globules and fat-needles, which are sufficiently characterised by their refracting property and their solubility in æther.
3. Elastic fibres and connective tissue.
4. Starch granules, to be recognised by their concentric arrangement

[* It is, however, doubtful if there is an emulsifying ferment.]

and by their property of staining blue with iodo-potassic-iodide solution. These bodies are frequently disintegrated and more or less dissolved by the process of digestion.

5. Vegetable cells of various forms.

In addition, the vomit in disease displays a great variety of fungoid growths (*W. de Bary*¹⁴¹), depending upon the nature of the underlying process. Amongst these :—

1. Mould-Fungi and scattered gonidia have occasionally been found. These are, so far as we know, devoid of pathological significance.

2. Yeasts.—(a) *Saccharomyces cerevisiae*. These are about the size

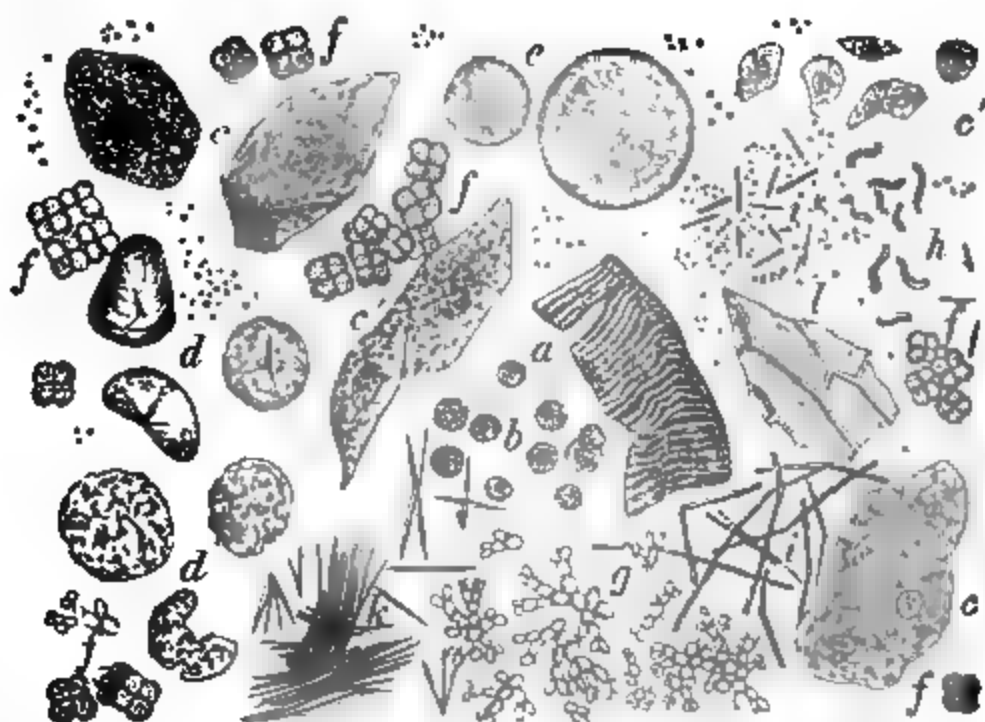


FIG. 67. Collective View of Vomited Matter (eye-piece III., objective 8a, Reichert).

- | | | |
|---|--|---|
| a. Muscle fibres. | e. Fat globules. | i. Various micro-organisms, such as bacilli and micrococci. |
| b. White blood-corpuscles. | f. Sarcina ventriculi. | j. Fat-needles; between them connective tissue derived from the food. |
| c, c'. Squamous epithelium. | g. Yeast-fungi. | k. Vegetable cells. |
| d. Starch grains, mostly already changed by the action of the digestive juices. | h. Forms resembling the <i>comma-bacillus</i> , found by the author once in the vomit of intestinal obstruction. | |

of leucocytes, and refract light powerfully. They cohere in groups of three or more, and stain deeply a brownish-yellow with iodine and iodide of potassium. Very often there are also to be seen elliptical bodies resembling *Saccharomyces ellipsoideus* (*Rees*).¹⁴²

(b.) Very small yeast-like fungi in thick clusters (fig. 67, g).

(c.) More seldom highly-refracting rod-like bodies of considerable length and thickness, which generally exhibit a single nucleus. They are rounded off at the ends, and are sometimes separate, sometimes strung together in strands. It would seem that these are the agents in lactic acid fermentation of sugar.

3. Fission-Fungi.—The forms belonging to this class are many and various.¹⁴³ They include rod-like bodies, which stain blue with iodo-potassic-iodide solution, bacilli and micrococci of every description, and in particular a bacillus which possesses the property of changing glycerine into alcohol by fermentation (fig. 67, *i*).

*Sarcinae ventriculi*¹⁴⁴ may easily be known by their shape, resembling that of wool-packs, their dark silver-grey tint, and their property of staining a deep mahogany-brown to reddish-brown with the iodo-potassic-iodide solution (fig. 67, *f*).

After this general view of its microscopical appearances, we shall advert to the physical, chemical, and microscopical characters of the vomit in certain diseases.

1. Acute Gastritis.—In this condition the vomit consists partly of mucus which has been swallowed, and partly of half-digested food residues. Microscopically it displays the appearances already detailed, which, however, are here subject to much variety, and notably a few red blood-corpuscles are generally to be seen.

Its chemical character varies greatly (*Ewald*).¹⁴⁵ At the outset of the affection, hydrochloric, and commonly lactic, acid in the free state are wanting. The addition of the former will establish a slow digestive process. *Ewald*¹⁴⁶ has been unable to determine the presence of fatty acids in notable quantity, and the proportion of pepsin would appear to be considerably diminished.

The vomit usually is coloured green from the admixture of bile pigment (biliverdin). It often also contains biliary acids. The first may be recognised by Gmelin's test (see chapter on *Urine*), and the latter by Pettenkofer's (see p. 92), or by means of the furfural and sulphuric acid reaction (see *Urine*). Much remains to be learnt as to the chemical peculiarity of the gastric contents in this condition.

2. Chronic Gastritis.—The vomit is a thin mucous fluid (*vomitus matutinus*), of alkaline, or it may be weakly acid, reaction. *Van der Velden* has shown that it always contains pepsin and hydrochloric acid, also organic acids, especially acetic and butyric acids. It is commonly rich in proteids, and notably peptones, which may be easily distinguished by the tests which will be subsequently described in connection with the examination of the urine. Bile pigment is also generally present.

Recent observations seem to show that it is possible to distinguish the different forms of gastric catarrh by the results of the chemical methods described above. Following the conclusions of *Ewald*, we may discriminate (1) Simple gastritis, (2) acid gastritis, (3) mucous catarrh, (4) atrophy.

1. In the first form, *simple gastritis*, the test breakfast is never followed by increased acidity; the proportion of hydrochloric acid is

diminished; the secretion contains little pepsin and milk-curdling ferment, and generally, though not always, includes lactic and fatty acids. On the addition of acid the secretion shows digestive activity.

2. In *acid gastritis* acidity is increased, especially that due to hydrochloric acid. In other respects the condition is that of simple gastritis.

3. In the third form, *mucous catarrh*, acidity is always slight and hydrochloric acid absent; there is abundance of propeptone, but no peptone. Milk-curdling ferment is absent, or it may develop only after a prolonged interval. Artificial digestion requires the addition of hydrochloric acid.

4. In *atrophic gastritis* the fasting stomach is usually empty, and its contents, after the administration of the test-meal, is free from mucus, and altogether wanting in pepsin, hydrochloric acid, and the milk-curdling ferment.¹⁴⁷

The observation of *Mathieu*¹⁴⁸ that mucus is indigestible has an important bearing on the study of these conditions. The fact noticed by *John*¹⁴⁹ that excessive acidity of the gastric juice impedes salivary digestion, and that acids, organic and inorganic, promote the secretion of saliva, is equally instructive. In the case of a gastro-duodenal catarrh complicating gastritis, the gastric juice exhibits the same variety in respect of the presence or absence of appreciable quantities of physiologically effective hydrochloric acid. The author has investigated three cases of this kind. In one the acid was altogether wanting, while in the other two it occurred in diminished proportion.

3. Chronic Ulcer of the Stomach.—The microscopical appearances of the vomit in this disease are those detailed under (2) in the last section. Otherwise it exhibits nothing distinctive. There can be no doubt that we have in the hyper-acidity which *Riegel's*¹⁵⁰ observations have connected with this condition in a large number of instances, a fact of the highest clinical significance. It may be estimated accurately by the method detailed in this chapter, or by titration. The proportion of hydrochloric acid in the stomach in a case of chronic gastric ulcer is, according to *Riegel*, 0.4–0.6 per cent., as against 0.1–0.2 per cent. in health.¹⁵¹ It must be mentioned, however, that the researches of *Ewald*, *Ritter* and *Hirsch*, and *Janczorowski*¹⁵² go to show that the increased acidity in connection with round ulcer of the stomach may undergo diminution with the further progress of the disease.

According to *Lenhartz*,¹⁵³ acid may be deficient in gastric ulcer. The digestion of carbohydrates is accomplished slowly. There is still need to verify these assertions by the application of trustworthy methods, such as those described in this chapter, and especially with reference to the precautions enjoined in a previous paragraph. The conflict-

ing statements quoted here must be in part referred to the want of such accurate observation.

The presence of *blood*, and its character, are facts of great significance.

1. When the haemorrhage is considerable, clots of blood are found, which are not at all, or but slightly changed.

2. More commonly the effused blood remains for a longer time in contact with the gastric juice, and is thereby altered in such a way that the oxyhaemoglobin is converted into haematin, and the vomit has the appearance of coffee-grounds.

When examined under the microscope in such a case, no blood-corpuscles whatever are to be seen in it, but in their place larger or smaller pigment masses. The blood may best be identified as such by *Teichmann's* haemin test and by the spectroscopic appearances of haematin. To obtain the latter, a portion of the vomit should be treated with caustic potash, filtered, and then examined with the spectroscope for the spectrum of haematin in alkaline solution (fig. 39).

It should be borne in mind that the exhibition of preparations of iron will impart to the vomit the same appearance as that due to blood; so also will the abundant partaking of red wine; and finally, the presence of bile pigment may cause it to assume a brownish-black colour.

Large quantities of blood (blood-pigments) may be found in the vomit in cases of duodenal ulcer with haemorrhage into the intestine.

4. Carcinoma of the Stomach.—The physical and microscopical character of the vomit in cases of cancer of the stomach are, in general, those of gastric ulcer. Sarcinæ in large quantities are a remarkably frequent manifestation. The blood is very seldom discharged unaltered, and it is usually represented only by colouring matter.

The chemical constitution of the gastric juice in cases of cancer has been made the subject of research by many observers, *van der Velden*, *Uffelmann*, *Ewald*,¹⁵⁴ and *Kredel*;¹⁵⁵ also by *v. Mering* and *Cahn*; and more especially by *Riegel*, *Korczynski*, and *Jaworski*.¹⁵⁶ Absence or diminution in quantity of hydrochloric acid is a point of special interest, and has been recently studied by many investigators.¹⁵⁷

The author has analysed the contents of the stomach in a great number of cases of cancer, and he has found that in many no trace of free hydrochloric acid could be shown by the colour-tests employed for the purpose. He has had seventy-six cases of cancer under observation during the last six years. Either the vomit or the gastric juice withdrawn after the administration of a test-meal—usually of milk or ham—was repeatedly examined. In sixty-one out of the seventy-six cases the reaction with Congo-paper, benzo-purpurin, and *Günzburg's* test, was either absent or very feebly evinced. The total acidity was also very low, ranging from 52–54, and the higher figure was obtained in only one

instance. In two cases, where the gastric juice was examined an hour after the administration of half a litre of milk, no result was obtained by the *Sjöqvist-Jaksch* method, while the same method applied to the gastric juice of a healthy person under like conditions showed 0.0301 grm. HCl in 100 cc. In three cases of cancer the acidity was remarkably high, viz., 90, 100, 126, and all the tests for free HCl gave decided results. The experiences of *O. Rosenbach* and *Waetzhold*¹⁵⁸ have been similar to these. The absence of hydrochloric acid, therefore, is by no means so constant as to warrant an absolute diagnosis on this ground alone. Moreover, in other conditions, such as amyloid degeneration of the gastric mucous membrane,¹⁵⁹ in stagnation of the contents of the stomach, in diabetes,¹⁶⁰ and in the febrile state, even without demonstrable disease of the stomach,¹⁶¹ the hydrochloric acid reactions may also fail. The author has determined their absence in two cases of carcinoma of the gall-bladder.

According to *Wolfram*,¹⁶² the gastric juice is devoid of HCl in the course of the infectious fevers, whereas in chronic febrile disorders it is of quite normal character, and *Leubuscher* and *Ziehen*¹⁶³ have shown that the acidity may be wanting in psychoses of various kinds.

In eighteen out of twenty-five cases of gastric cancer investigated by the methods given in this chapter, the efficiency of the acid was proved. The author is therefore unable to endorse *Riegel's* statement. *Boas'*¹⁶⁴ observations seem to show that the presence of large amounts of lactic acid are a symptom of cancer of the stomach, and this is confirmed by others.¹⁶⁵ *Rosenheim*¹⁶⁶ also has asserted that in the great majority of cases of this disease the perchloride of iron reaction is very pronounced. The process for the detection of lactic acid by *Boas'* method is given in this chapter. *Oppler*¹⁶⁷ and others have found in the cancerous stomach elongated unsegmented bacilli, which effect the decomposition of glucose with lactic acid as a product, and these have been thought to indicate the existence of cancer. *G. Klempner*¹⁶⁸ regards the presence of lactic acid simply as the result of prolonged stagnation of the stomach contents.

The evidence as to the diagnostic significance of lactic acid at present at the author's disposal is the following. In twenty-nine cases clinically diagnosed as cancer of the stomach and investigated in his clinic between the years 1892-95, lactic acid was always found in the gastric contents. This fact would have greater weight were it not that in one case diagnosed as cancer of the pylorus, but which the autopsy showed to be gastric ulcer, the stomach had also yielded abundance of the acid. In three other cases, also of gastric ulcer, the same was found; in one of cancer of the pancreas also; and in another of uncertain nature, where the recovery of the patient excluded the possibility of its being cancer.

In this case microscopical examination of the stomach contents showed disintegration of the gastric mucous membrane, pus cells, &c.—the indications of phlegmonous ulceration of the organ. The patient, a man, was the subject of a remarkable aberration of the intellect.

From what has been said, it follows that lactic acid occurs very frequently in cancer of the stomach, but that it occurs in other conditions as well, and is not, therefore, pathognomonic of cancer, but is rather to be taken as indicating retention and decomposition within the stomach. Nevertheless, this sign, taken together with others, as the absence of hydrochloric acid, is of great value in the recognition of cancer. It is necessary to add that an inference cannot safely be based on the presence of lactic acid when this has been determined *only* by *Uffelmann's* unsatisfactory test. Taken, however, in conjunction with the other clinical symptoms of cancer, we have in this circumstance important evidence of the disease. *Riegel* points out the important fact that the gastric juice in this affection has entirely lost the digestive property. Concerning the secretion of *pepsin* in carcinoma, it would appear that this, as well as the milk-curdling ferment, are secreted to the end. The tests for pepsin are described above, and its quantitative estimation may be effected by *E. Schütz's* method.

5. Dilatation of the Stomach.—The character of the gastric contents in this condition is subject to variety, according to the cause of the dilatation. Nevertheless, there are certain general features which belong to all cases, and these will be considered first. The remnants of undigested food are visible many hours after a meal. Microscopically there is a profusion of micro-organisms of all kinds, and yeast-forms are rarely absent. Chemical analysis usually discloses an excessive proportion of volatile fatty acids and of lactic acid.

In dilatation from chronic gastritis the contents partake of the character of the latter disease.

When the primary condition is pyloric stenosis, with gastric ulcer, physiologically active HCl is present in great excess. In one case the author found that the unfiltered gastric juice withdrawn in the morning before food had been taken contained 0.4629 grm. HCl to 100 cc.; and when the stomach had been thoroughly washed out, and milk administered, the gastric juice examined half-an-hour afterwards contained 0.1374 grm.

In dilatation from cancer of the pylorus free HCl is either absent or diminished in quantity. This is true also of dilatation with atrophy of the stomach, but the rule is not without exceptions.

6. Parasitic Affections of the Stomach.—(a.) In but one instance as yet—a case of favus—have the characteristic appearances of this condition been detected in the stomach (*Kundrat*).¹⁰⁹

(b.) Extensive patches of thrush are sometimes formed in the stomach, and in such cases the vomit contains masses of the thrush fungus.

7. Diphtheria.—It very rarely happens that a diphtheritic condition of the mucous membrane extends from the pharynx as far as the stomach. When it is so, the vomit exhibits the appearances described in Chapter ii. Croupous formations which are not diphtheritic sometimes occur.

8. Fæcal Substances in the Vomit.—Formed masses of fæces are never discharged by the mouth; but in cases of occlusion or partial paralysis of the intestine, its contents may become mingled with those of the stomach, and brought up with the vomit, which then has an intensely fæculent odour, a yellowish-green colour, and a feebly acid or alkaline reaction. When derived chiefly from the small intestines, the vomited matter will contain bile acids and pigment and abundance of fat. These may be detected by chemical examination. Microscopically it shows nothing distinctive; but on one occasion the author found in such a discharge a quantity of large fungi which closely resembled the comma-bacillus (fig. 73).

9. Pus.—In rare cases pus occurs in the vomit. It indicates suppuration in the walls of the stomach, or the rupture into it of an abscess, from some neighbouring viscus.

10. Animal Parasites.—Amongst the Entozoa, *Ascaris lumbricoides*, *Oxyuris vermicularis*, and *Anchylostoma duodenale* have been obtained from the stomach. Other worms, such as *Trichina*, are exceptional manifestations, and the hooklets of *Echinococcus* and hydatid cysts are occasionally present. *Gerhardt*¹⁷⁰ has found dipteral larvæ in the secretion, where they give rise to the symptoms of gastritis. Similar observations have been made by *Senator*,¹⁷¹ *Hildebrandt*,¹⁷² and *Finlayson*.

11. Constitution of the Vomit in Poisoning.¹⁷³

1. Poisoning with Acids.—In all cases of poisoning with strong mineral or organic acids, the vomit acquires a powerfully acid reaction. It displays a blackened mass of altered blood and charred tissues in cases where the quantity of the poison has been sufficient to cause these effects. The appearance presented in all cases of poisoning with strong acids is the same. To distinguish one from the other in the contents of the stomach, the sense of smell will serve in some instances, as, e.g., for acetic acid, while in others resort must be had to the methods of analytical chemistry. It is important in certain cases to remember that the vomit may contain an abnormal proportion of organic and inorganic (hydrochloric and lactic) acids, altogether independently of poisoning.

(a.) Detection of Sulphuric Acid.—The presence of sulphuric acid may be ascertained thus:—The vomit is mixed with a large bulk of

distilled water, and put aside for several hours, during which it is frequently stirred. It is then filtered, and the precipitate repeatedly washed with water on the filter. The filtrate is next collected, and evaporated on the water-bath until the fluid begins to blacken. It is then allowed to cool, mixed with twice its bulk of alcohol, and, after standing for some hours, again filtered. The filtrate, diluted with water, is once more evaporated on the water-bath until the alcohol is entirely driven off. The fluid remaining may then be tested for sulphuric acid. The addition of chloride of barium solution or of lead nitrate should give a white precipitate, showing the presence of sulphuric acid or its salts.

(b.) **Detection of Nitric Acid.**—The vomit, which is generally stained somewhat of a yellow hue from the formation of xanthoproteic acid, is mixed with water, boiled, and filtered. The reaction of the filtrate is tested, and, if found to be acid, it is neutralised with caustic potash, and evaporated to a small bulk. When allowed to cool, it should deposit crystals of nitrate of potash, which will give the following reactions:—

1. To a solution of the crystals concentrated sulphuric acid is added, and when the mixture is quite cool, a little sulphate of iron solution is poured upon its surface. At the point of contact of the two fluids a deep-brown zone shows the presence of nitric acid. The test is applicable only *where the brown coloration is not obtained with sulphuric acid alone.*

2. A solution of brucin in sulphuric acid is placed in a test-tube, and a little of the fluid supposed to contain nitric acid is poured on its surface. At the point of contact of the two a red coloration takes place if nitric acid is present.

The modes of testing for **hydrochloric acid** have already been given.

(c.) **Oxalic Acid.**—The contents of the stomach are partially evaporated and extracted with alcohol, the alcohol evaporated, and the residue, dissolved in water, is treated with acetic acid and solution of chloride of calcium. A precipitate of oxalate of calcium forms, and the characteristic crystals may be distinguished by the microscope.

2. Poisoning with Alkalies.—The vomit in such cases is a viscid, glistening, and strongly alkaline fluid. Where a concentrated solution of a caustic alkali has been taken, the ejected tissues are charred and brown, as in the case of acids. The determination chemically of the character of the poison is often attended with great difficulty, and in other instances, again, it is very easy.

Where *ammonia* has been taken, the vomit, if examined immediately after the poisoning, will emit the characteristic odour of that body, and further evidence of its presence may be obtained by holding over the fluid a glass rod moistened with hydrochloric acid, when fumes of sal ammoniac will be given off. The detection of caustic potash and of

caustic soda, on the other hand, involves much difficulty on account of the readiness with which they are converted into their carbonates.

The process for the detection of *chlorate of potash* in the vomit demands a special notice. *E. Ludwig's*¹⁷⁴ method is as follows:—The vomit, if not already acid, is rendered slightly so by the addition of acetic acid, heated and maintained for one minute at the boiling-point, filtered, the filtrate evaporated to a small bulk on the water-bath, and allowed to settle undisturbed. The salt then separates in the form of crystals, which are dried between folds of blotting-paper and tested in the following manner:—

1. They are treated with dilute hydrochloric acid and warmed; the fluid assumes a greenish-yellow colour, and chlorine gas is evolved. The reaction may be obtained at ordinary temperatures by the addition of strong hydrochloric acid.

2. The crystals are dissolved in water, or, where none have been deposited, the liquid is evaporated and a solution of indigo and dilute sulphuric acid is added. If chlorate of potash be present, on the further addition of a watery solution of sulphurous acid or hyposulphite of soda, the fluid changes from blue to yellow, or altogether loses its colour.

3. Poisoning with Metals and Metalloids. — (a.) **Poisoning with Salts of Lead.** — After the lapse of a few hours, a quantity of grey or blackish-grey substance is vomited. For the detection in this of compounds of lead, it should be partially evaporated on the water-bath, and organic matter decomposed by treatment with reagents.

For this purpose *E. Ludwig* recommends the process of *Fresenius* and *Babo*:—The vomit is placed in a large porcelain dish, mixed with about its own weight of a 20 per cent. solution of hydrochloric acid, and 3 to 5 grms. of chlorate of potash added, when the vessel is covered and allowed to stand for about twelve hours. The mixture is then heated to 60° in the water-bath. When the evolution of gas has ceased, more chlorate of potash is added to the brown mass, and this process is continued until the fluid ceases to form a brown colour. Should the fluid become unduly concentrated, it should be further diluted with water. If in this way the organic material is not entirely decomposed, hydrochloric acid must be again added, and the requisite quantity of chlorate of potash supplied as before. The mixture is then evaporated on the water-bath until the odour of chlorine is no longer perceived, when it is diluted with twice its bulk of water, passed through a filter which has been moistened with water, washed with a large quantity of water, and the washings finally collected and added to the filtrate. To the fluid so obtained sulphuretted hydrogen is added to saturation. The resulting dark precipitate is then filtered off, washed with sulphuretted hydrogen water, dried, and dissolved in nitric acid in the following

manner :—It is placed in a porcelain dish, and pure (chlorine free) nitric acid is added in drops until the whole has the consistence of a thin fluid, when it is evaporated to dryness on the water-bath, and the residue dissolved in boiling water and filtered. A white insoluble residue of sulphate of lead may remain. This may be reduced to metallic lead by the addition of soda and combustion on charcoal in the reducing zone of a blow-pipe flame.

The addition of sulphuric acid will cause the formation of a *white* precipitate of sulphate of lead, and chromate of potash will give a *yellow* precipitate if lead be present.

The quantitative estimation may be effected in the same way.

Salts of lead in the vomit may be detected by another very simple process :—A strip of magnesium, free from lead, is placed in the fluid, when metallic lead will be deposited upon it, and can then be dissolved in nitric acid, and subsequently proceeded with as above.

(b.) **Poisoning with Salts of Mercury.**—Poisoning with compounds of mercury is very often attended with vomiting. The vomit in such cases differs greatly according to the strength of the poison. When large quantities of corrosive sublimate have been taken, pain is experienced in the region of the stomach, and shreds of tissue stained brown with haematin are apt to be discharged in the vomit.

The salts of mercury may be detected in the vomit in the same manner as the compounds of lead. Sulphide of mercury is formed in the process, and from this the metal is obtained thus :—To the precipitate carbonate of soda and cyanide of potassium are added, the mixture dried, placed in a test-tube and heated, when the metal is sublimed in the upper part of the tube.

The presence of mercury in the vomit may be shown directly as follows :—Granulated zinc (*E. Ludwig*)¹⁷⁵ or brass wire (*Fürbringer*)¹⁷⁶ is placed in the substances to be examined, which have been previously acidulated with hydrochloric acid; the mixture heated for an hour in the water-bath, and exhausted first with water and then with alcohol, and finally washed with æther and allowed to dry in the air. The brass wire is then placed in a test-tube and heated, when the metal is deposited on the sides of the tube. If now, while the test-tube and its contents are still hot, a small piece of iodine be introduced, the vapour of iodine given off will change metallic mercury into mercuric iodide with the development of a beautiful red colour (*Schneider*).¹⁷⁷ In a similar manner, mercury obtained by other methods may be converted into mercuric iodide. If, at the same time, the vomited matter contains an abundance of organic substances, it will be necessary to remove the latter by *Fresenius* and *Babo's* method before introducing the granulated zinc or brass-foil.

It should be mentioned that in the distillation of the suspected vomit, metallic mercury may pass over with the watery vapour (*Lecco*¹⁷⁸), and it may also form from the reduction of corrosive sublimate in the process of distillation.

(c.) **Poisoning with Salts of Copper.**—When sulphate of copper has been taken, it imparts a greenish-blue tint to the vomit. In poisoning with the copper salts of acetic acid (verdigris, the commonest form), the green colour may be present, or there may be nothing distinctive in the appearance of the gastric contents. The recognition of the poison may be effected by the process described under (a.) *Poisoning with Lead Salts*. The sulphide of copper so obtained is dissolved in nitric acid, when the presence of the metal will be shown by the blue colour of the solution, which becomes deeper on the addition of ammonia. If the latter reagent causes a precipitate to fall, the fluid is filtered and the filtrate acidulated with hydrochloric acid. A portion of the filtrate is treated with yellow prussiate of potash, when a reddish-brown precipitate forms. In another portion a piece of iron-foil is placed, and after a little while the metallic copper present is deposited on its surface as a red coating.

It must not be forgotten that traces of copper occur in every organ.

(d.) **Arsenic Poisoning.**—The administration of large doses of arsenious acid, Fowler's solution, or of certain mineral waters abounding in arsenic (such as those of Roncegno and Levico, &c.), is followed after a short interval by the vomiting of a copious fluid deeply stained with bile. Where arsenious acid (white arsenic) is the poison in question, a careful naked-eye and microscopical examination of the vomit will often afford accurate information as to its nature. In such a case larger or smaller particles of this substance are usually to be seen. When these white particles are removed with forceps, freed from other impurities by repeated cleansing, washed with cold water, and dissolved in a test-tube containing a small quantity of boiling water and then allowed to cool, crystals of arsenious acid separate, and may be recognised microscopically as small octahedral forms. When the crystals are heated with soda on a piece of carbon in the reducing zone of the blow-pipe flame, the characteristic odour of garlic is evolved, and if a specimen be heated with carbon in a test-tube, a metallic deposit forms in the upper cool part of the tube.

A more accurate method is to remove organic substances with chlorate of potash and hydrochloric acid, then to heat the remaining fluid for a long time at 60° with sulphuretted hydrogen. A yellow precipitate of sulphide of arsenic forms. This is dissolved in sulphide of ammonium, filtered, the filtrate evaporated to dryness, allowed to cool, some drops of concentrated nitric acid added, heated with more nitric

acid until the evolution of gas has ceased and reddish-brown fumes are no longer given off. The fluid is then concentrated to a small bulk in the water-bath, diluted with a little water, and treated with sodium carbonate until its reaction is distinctly alkaline. It is then evaporated to dryness on the water-bath; the dried residue is fused with a quantity of carbonate and nitrate of sodium, allowed to cool, several times exhausted with water, and filtered. The filtrate is repeatedly treated with small quantities of dilute sulphuric acid until effervescence has ceased, when more sulphuric acid is added, and the fluid evaporated first on the water-bath and afterwards over a flame until white fumes are given off. The residue is then allowed to cool and is dissolved in cold water. Zinc and sulphuric acid, both free from arsenic, are placed in an apparatus for generating hydrogen.¹⁷⁹ The hydrogen set free is purified and dried by being passed through a tube which is fitted to the apparatus, and contains solid caustic potash and granular chloride of calcium. To the first tube is adapted, by an air-tight connection, another tube constricted in two or three places and terminating in a point. When all the air has been driven out, the hydrogen escaping from the point of the terminal tube is ignited, and then the fluid to be tested is poured into the apparatus. The tube is next heated in front of the place where its calibre begins to diminish, and if arseniuretted hydrogen is mixed with the hydrogen gas evolved, metallic arsenic is deposited in the constricted portion.

A further test is as follows:—The flame is extinguished and the gas conducted into a solution of nitrate of silver. A blackish-grey precipitate of metallic silver separates, and the filtrate, on the careful addition of ammonia, will give a further yellow deposit of arsenite of silver.

(e.) **Phosphorus Poisoning.**—Poisoning with phosphorus is always attended with persistent and severe vomiting, which may last for whole days. The discharged substances are free from blood, shreds of tissue, and the other signs of a formidable organic lesion. When solid phosphorus in large quantity has been taken, the gastric contents emit the characteristic odour of phosphorus and are luminous in the dark. It should be noted, however, that these properties are lost in presence of alcohol, oil of turpentine, and chloroform.

For the detection of phosphorus, the vomit is distilled with sulphuric acid in the dark, and the retort connected with a glass condenser (*Mitscherlich*). The presence of the poison is shown by the appearance of luminous rings where the phosphorus fumes come in contact with the cold water.

Scherer has also devised an admirable method for this purpose. The vomit is enclosed in a flask provided with an air-tight stopper, and two test-papers—one saturated with nitrate of silver and the other with

acetate of lead—are placed in it. If phosphorus be present, the first of these will be blackened, whilst the other remains unchanged.¹⁸⁰

4. Poisoning with Alkaloids.*—(a.) Morphia Poisoning.—The earlier stages of poisoning with morphia are generally attended with vomiting, and in all cases where it has been taken by the mouth the alkaloid can be detected in the ejected contents of the stomach. *Alt* and *Hitzig*¹⁸¹ have shown that, even when it has been injected subcutaneously, the drug will be found in the stomach about an hour afterwards. In such a case, therefore, the stomach should be washed out, and the washings tested for morphia. The *Stas-Otto* method¹⁸² for its separation may be employed. In this, the vomit is placed in a flask, and digested with alcohol and tartaric acid on the water-bath, allowed to cool, and filtered; the alcoholic extract is heated on the water-bath at a moderate temperature (60°) until the spirit is entirely driven off, when the remaining watery solution is filtered. The new filtrate is evaporated to a syrupy consistence on the water-bath, and the residue extracted with alcohol. In doing this, the latter should be cautiously added, little by little, until a flocculent precipitate forms, and then in greater quantities until no further turbidity occurs. The alcoholic solution is then filtered, and the filtrate evaporated on the water-bath and dissolved in a little water. The still acid watery solution is next shaken up with æther, in order to eliminate other alkaloids and resinous substances, after which it is rendered alkaline with caustic potash, and again shaken up with æther. Any nicotin and atropin present (see below) are dissolved in this way. The residue is treated with sal-ammoniac and repeatedly extracted with warm amylic alcohol, which takes up morphia. The amylic alcohol extract is next collected, filtered, and evaporated to dryness on the water-bath. The residue is dissolved in acidulated water, which is repeatedly added for the purpose. It is then filtered, extracted with amylic alcohol, neutralised with ammonia, and again extracted with warm amylic alcohol, which is driven off by evaporation. Finally, the residue may be tested thus:—

1. To one portion a freshly prepared solution of molybdate of soda and concentrated sulphuric acid (1 cc. sulphuric acid and 5-10 grms. of molybdate of soda—*Fröhde's reagent*) is added. If morphia be present, the fluid turns first violet, and, changing through blue and green, becomes finally a pale red.

2. Another portion is dissolved in water acidulated with hydrochloric acid, evaporated on the water-bath to dryness, and treated with a few

* In this connection we shall consider only such alkaloids as are most frequently the subject of investigation by the physician. For more detailed information on the entire subject, the text-books of *F. C. Schneider*, *J. Otto*, *Kobert*, and *E. Ludwig* may be consulted; [and in English those of *A. Wynter Blyth* and *Dixon Mann*].

drops of a very dilute solution of perchloride of iron, which should be *free from hydrochloric acid*. A blue colour shows the presence of morphia.

An acid-free solution of perchloride of iron may be best prepared by dissolving the sublimated salt in water.

(b.) **Poisoning with Nicotin.**—Vomiting frequently occurs in this condition, and the poison may be separated from the gastric contents by the *Stas-Otto* method. The alkaloid is extracted with æther from the alkaline solution of the evaporation residue (*vide supra*), and when the æther is driven off at a low temperature (30° C.) on the water-bath, nicotin remains as a brown or yellow mass. If this be dissolved in æther and an æthereal solution of iodine added, an oily substance forms, from which ruby-red needles (*Roussin's crystals*) slowly separate.

(c.) **Poisoning with Atropin.**—In atropin-poisoning with the pure alkaloid, whether taken by the mouth or administered subcutaneously, vomiting rarely occurs ; but it is a frequent event in poisoning with the berries of the deadly nightshade. Under such circumstances, the condition is usually rendered evident by the discovery of the berries in the ejected substances, together with the clinical symptoms (mydriasis, &c.). If there should be any doubt as to the character of the toxic agent, recourse may be had to the *Stas-Otto* method, when the alkaloid will be taken up by æther from an alkaline solution of the residue, and may be recognised by the following tests. The æther being driven off :—

1. A portion of the residue dissolved in water, to which a trace of acid has been added, is dropped upon the conjunctiva of an animal (cat or rabbit). After the lapse of from 6-20 minutes paralysis of the sphincter fibres of the iris ensues, and the pupils are widely dilated : 0.01 mgrm. of atropin suffices to produce this effect.

2. A specimen of the residue is dissolved in a few drops of fuming nitric acid, and the solution evaporated on the water-bath : a colourless substance remains, and this, when allowed to cool, and subsequently treated with alcoholic solution of caustic potash, turns first violet, and afterwards a cherry-red colour.

(d.) **Poisoning with Ptomaines and Toxalbumins.**¹⁸³—The ingestion of putrid flesh is occasionally attended with severe symptoms of poisoning, and there can be no doubt that in many cases of so-called acute gastritis, where the use of certain articles of diet—such as liver, kidneys, and oysters—has been followed by nausea, vomiting, and profuse diarrhoea and a retarded pulse, the true cause is to be found in the toxic effects of ptomaines and toxalbumins. To the same category may be referred the conditions known as ammoniæmia (retention-toxicosis, *v. Jaksch*,¹⁸⁴ see *Urine*), and the cerebral symptoms (coma carcinoma-

tosum, *v. Jaksch*¹⁸⁵), which sometimes develop in the course of cancer.¹⁸⁶ The poisonous substances are probably diamines and toxalbumins, which are known to be formed in these processes. Further research is needed for the elucidation of this subject, and in questionable cases the vomit should be tested for ptomaines. In doing this, the observer should be guarded in his conclusions, since peptone (which is normally a constituent of the gastric contents) is known to yield toxic substances of an alkaloid nature.

In view of the great importance which the study of these bodies has acquired of late years, and also because it enters into the pathology of other secretions besides that now under consideration, the subject is dealt with here at some length. The references, which are omitted, will be found in *Brieger* and the other authorities quoted.

The *Stas-Otto* method may be employed for the separation of ptomaines from the vomit. Such of these bodies, however, as have yet been distinguished exhibit a very remarkable variety in respect of their chemical character. Some may be derived by extraction with æther from an acid, others from an alkaline medium, whilst a third class is distinguished by the fact that its members are soluble only in amylic alcohol, chloroform, or benzol. Others, again, are insoluble in amylic alcohol. It will be seen from this that the application of the *Stas-Otto* process must be supplemented by extracting the derived substances with various media, and even then it will sometimes happen that the effort is attended with failure.

For such cases *Brieger's* method¹⁸⁷ may be employed. This is briefly as follows :—If the material to be examined contains solid substances, these are first reduced to small particles; sufficient hydrochloric acid is then added to render the whole feebly acid, and the mixture boiled for a few minutes and filtered. The filtrate is evaporated, at first over a flame, and subsequently on the water-bath, until it has attained a syrupy consistence. It may be noted, however, that in view of the instability of the bodies sought, it is advisable to evaporate in vacuum, and at the lowest possible temperature (*r. Jaksch*)—a precaution which, as *Brieger*¹⁸⁸ suggests, should also be taken where foul-smelling substances are the subject of manipulation. The thick fluid is mixed with 96 per cent. alcohol (filtered), and the filtrate treated with warm alcoholic solution of acetate of lead. The lead precipitate which forms is now filtered off, and the filtrate concentrated—here again preferably in vacuum—to the same consistence as before, when it is again taken up in 96 per cent. alcohol. The alcohol is driven off by evaporation, and the residue, dissolved in water, is freed from lead by the addition of sulphuretted hydrogen and filtering. The filtrate is acidulated with a little dilute hydrochloric acid, and concentrated (in vacuum) to the consistence of

syrup. It is then diluted with alcohol, and alcoholic solution of mercuric chloride added. The resulting precipitate is boiled in water, and certain ptomaines may separate at this stage in consequence of the different solubilities of the double salts of mercury. The better to secure this, the precipitate may be treated successively with water at various temperatures. Should it be thought that the lead precipitate may have retained some of the ptomaines, it may be suspended in water, the lead converted into its sulphide, and the fluid treated in the manner just described.

The solution obtained as above is filtered, and the filtrate, already freed from alcohol and mercury, is evaporated, the hydrochloric acid all but completely neutralised with sodium carbonate, and the residue again extracted with alcohol, after which it is dissolved in water, neutralised with soda, again acidulated with nitric acid, and precipitated with phosphomolybdic acid. The double phosphomolybdate is filtered off and decomposed by neutral acetate of lead—an object attained more readily by the application of heat on the water-bath. The lead is then removed by means of sulphuretted hydrogen, and the fluid concentrated and treated with alcohol. Several ptomaines are thus separated as hydrochlorates, and may be obtained in the form of double salts of gold or platinic chloride and of picric acid. From these the hydrochlorates are again derived by precipitation with sulphuretted hydrogen; and in the case of the picric acid compounds, extracting with water, acidulating with hydrochloric acid, and finally removing the picric acid by shaking it up with æther.

The next step is to ascertain if any ptomaines remain in the phosphomolybdic acid filtrate after the precipitation of phosphomolybdic acid.

The above is a mere sketch of the process, which, moreover, needs to be modified in many instances.

For the detection of the basic compounds which occur in the secretions as diamines, the best method is that of *Baumann* and *v. Udransky*, in which these bodies are converted into their benzoic compounds by the action of benzoyl chloride and caustic potash. By this means these observers discovered cadaverin (pentamethylendiamine) in the urine, and established the identity of *Brieger's* putrescin with tetramethylendiamine¹⁸⁹ (see chapter on the *Urine*).

Ptomaines (animal alkaloids) yield the characteristic reactions of alkaloids; but otherwise they are not known to possess any distinctive chemical or physiological properties.¹⁹⁰

The characteristic reactions which are common to all the alkaloids are the following (*Otto, E. Ludwig*).¹⁹¹ With—

1. *Iodine and iodide of potassium solution*, a brown flocculent precipitate; most

readily obtained from the solution of the alkaloid which has been acidulated with sulphuric acid.

2. *Mercuric and potassium iodide*, a white or yellow precipitate, insoluble in water and dilute acid.

3. *The iodide of bismuth and potash*, an orange precipitate in a solution acidulated with dilute sulphuric acid.

4. *Phosphomolybdic acid*, a bright or brownish yellow precipitate, insoluble in water and dilute mineral acids.

5. *Metatungstic and phosphotungstic acids*, a white flocculent precipitate, with difficulty soluble in water and dilute acid. (This, according to *E. Ludwig*, affords a particularly sensitive test.)

6. *Tannin*, in neutral or feebly acid solutions, a yellow or white precipitate.

7. *Platinic chloride*, a whitish-yellow or citron precipitate, which is sometimes readily soluble in water, and but slightly so in alcohol.

8. *Chloride of gold*, a yellow or whitish-yellow precipitate, which may be amorphous or crystalline.

The animal alkaloids already distinguished as occurring in the human system are sufficiently numerous. They have been detected in the faeces, the urine, and the milk, and they will find appropriate notice under the separate headings in this work. They are known by their effects both in health and disease, and, in many instances, have been separated as definite bodies from the secretions. Such products result from the decomposition of articles of food. Thus *Vaughan*¹⁹² obtained one of these bodies (tyrotoxin) from rotten cheese and bad milk, and he supposes that the substance in question is diazobenzol. *Ehrenberg*¹⁹³ found a similar body in putrid sausages. Special mention should be made also of ptomaine-atropin, a basic compound, which has been discovered in the latter food. In all cases where the clinical symptoms are those of poisoning, and include severe vomiting, the methods described above may serve to elucidate the matter; but it should not be forgotten, meanwhile, that the presence of peptone may be a source of ambiguity, since it may yield similar poisons. If the vomit or secretion is to be tested for toxalbumins, *Brieger* and *Fränkel's* process¹⁹⁴ may be used.

5. Poisoning with Ethyl Alcohol.—In acute poisoning with alcohol (ethylic alcohol), the vomit emits the characteristic odour of that substance. For its more accurate recognition, the gastric contents are first diluted with water, and, if very acid, carefully neutralised with caustic potash, and then distilled by steam.

The distillate may then be submitted to the following tests:—

1. To a portion a few drops of benzoyl chloride and a little caustic potash are added. When the mixture is heated and again allowed to cool, the presence of alcohol is shown by the characteristic odour of benzoylethylic æther (*Berthelet*).¹⁹⁵

2. With a small portion an equal volume of concentrated sulphuric

acid is cautiously mixed, a little powdered sodium acetate added, and the mixture heated. The characteristic odour of acetic æther shows the presence of alcohol (*Otto, E. Ludwig*).¹⁹⁶

6. Poisoning with Chloroform.—Chloroform may be detected either directly in the vomit or in the fluid obtained from its distillation. In either case, the following reactions will disclose its presence:—

1. A little thymol dissolved in caustic potash is added to the fluid to be tested, and the latter is then heated. If chloroform be present, the preparation assumes a dark violet tint (*Vitali*)¹⁹⁷, and if β -naphthol be used instead of thymol, a blue colour results (*Lustgarten*).¹⁹⁸

2. A few drops of alcoholic solution of caustic potash and a little aniline are heated with the distillate of the gastric contents. If chloroform be present, isocyanophenyl is formed, and may readily be recognised by its unpleasant odour (*Hoffmann*).

In a case of poisoning with chloroform, where the drug was taken by the mouth, the author was unable to find any trace of it in the substances vomited three hours afterwards, although the toxic symptoms were well marked.

7. Poisoning with Carbolic Acid.—When a poisonous dose of carbolic acid has been administered by the mouth, the vomit usually emits the characteristic odour of that substance. The presence of carbolic acid in the ejected substances may be ascertained directly by the following tests:—

1. Bromine water yields with a fluid containing carbolic acid a yellow crystalline precipitate of tribromophenol.

2. A solution of perchloride of iron colours dark violet in presence of carbolic acid.

In testing for these reactions, it is well previously to filter the vomit, if necessary washing it upon the filter with water. If the reactions are not obtained in this way, the filtrate is distilled with a little sulphuric acid, and both tests are again applied to the distillate.¹⁹⁹

It is to be borne in mind that carbolic acid may form in considerable quantity in the intestine in certain morbid states, and may then, as in intestinal obstruction, find its way into the vomit. (See chapter on the *Urine*.)

8. Poisoning with Nitro-Benzol and Aniline.—(a.) *Nitro-Benzol*.—The presence of nitro-benzol in the vomit may be detected frequently by the characteristic odour, which resembles that of the oil of bitter almonds. To separate this substance from the gastric contents, the latter is distilled with a little sulphuric acid. The distillate will contain oily drops which are soluble in æther. Aniline may be derived from nitro-benzol by the addition of granular zinc and dilute hydrochloric acid. When reduction is effected in this way, the fluid is ren-

dered alkaline with caustic potash, and the aniline formed is extracted with æther. The oily residue after the expulsion of the æther will yield the following reactions:—

1. If the aniline solution be treated with hydrochloric acid, and a shaving of pine-wood be placed in it, the latter assumes a deep yellow colour.

2. A drop of oil is suspended in water, and a few drops of dilute solution of chloride of lime or of a very dilute solution of sulphide of ammonium added : the fluid assumes a rosy-red colour (*Jacquemin*).²⁰⁰

3. A very sensitive test is that of *E. Ludwig*.²⁰¹—A watery solution of aniline colours a dark blue on the addition of a watery solution of carbolic acid and hypochlorite of soda ; and this colour changes to red on the further addition of hydrochloric acid.

4. The isocyanphenyl test serves well for the detection of aniline formed in the decomposition of nitro-benzol. It was introduced by *A. Flückiger*²⁰² as a test for acetanilide (antifebrin). To the fluid under examination a few drops of caustic potash and chloroform are added. It is then shaken up and heated, and, when again allowed to cool, should emit the disagreeable odour of isocyanphenyl.

(b.) *Aniline*.—Poisoning with aniline also is often attended with vomiting. The vomit is diluted with water and distilled with a little sulphuric acid. The distillate is extracted with æther, and when the latter has been driven off by evaporation an oily substance remains, to which the above tests (1 to 4) for nitro-benzol may be applied.

9. Poisoning with Prussic Acid.—This condition may usually be recognised by the characteristic odour of oil of bitter almonds. For the detection of the poison the vomit is treated with a small quantity of tartaric acid, and distilled. Hydrocyanic acid passes over with the distillate. From its presence, however, poisoning with the drug is to be inferred only when it can be ascertained that the vomit is free from innocuous double salts of cyanogen, as, for instance, the yellow or red prussiate of potash.

The best test is to add solution of ferric chloride and sulphate of iron to a little of the filtered fluid. With this last reagent yellow ferrocyanide of potassium gives a white precipitate, which soon changes to a bright blue, while with the perchloride of iron it forms a Prussian-blue precipitate. Red ferricyanide of potassium gives a dark blue precipitate with sulphate of iron, and a deep brown coloration with the perchloride. Should both the above-mentioned substances be present, the following process may be applied:²⁰³—The fluid is acidulated with sulphuric acid, and treated with an excess of carbonate of lime. The corresponding lime-salts take the place of ferro- or ferri-cyanide of potassium, and only so much hydrocyanic acid as is not combined in

the double salts of cyanogen passes over with the distillate. The latter is then tested for hydrocyanic acid in the following manner :—

1. To a few cc. rendered alkaline with caustic potash, a few drops of a freshly-prepared solution of cupric sulphate are added ; the mixture is heated, maintained for the space of one minute at boiling-point (*Ludwig*), and allowed to cool. It is then rendered strongly acid with hydrochloric acid. A blue coloration of the fluid results, and when allowed to stand for some time, a flocculent Prussian-blue precipitate settles in it.

2. To a few drops of the distillate a yellow (holding poly-sulphide of ammonium) solution of sulphide of ammonium is added, and the mixture boiled until the yellow colour entirely disappears. It is then allowed to cool, and perchloride of iron and hydrochloric acid added. If hydrocyanic acid be present, the preparation assumes a red colour (sulphocyanide of iron). *E. Ludwig*²⁰⁴ applies this test in a different manner :—The fluid is treated with yellow solution of sulphide of ammonium in excess, a few drops of caustic potash added, and the mixture evaporated to dryness. The residue is exhausted with water, treated with hydrochloric acid, and filtered. The filtrate is then tested with solution of perchloride of iron, when a blood-red colour should develop.

3. Another very admirable test has lately been suggested by *Vortmann*.²⁰⁵ To the fluid to be tested a few drops of nitrite of potash are added, and then two to four drops of the solution of perchloride of iron, and finally dilute sulphuric acid, until the yellowish-brown colour of the basic ferric salt formed in the beginning of the reaction has changed into a light yellow. The solution is then heated to boiling, allowed to cool, treated with ammonia and filtered, and to the filtrate a small quantity of a colourless solution of sulphide of ammonium is added. The presence of a minute quantity of hydrocyanic acid will be shown by a bluish-green, of large quantities by a beautiful violet-red colour. *Vortmann* has named this the “nitro-prusside test.”²⁰⁶

The vomit induced by a number of other poisons, as carbonic oxide gas, sulphuretted hydrogen, &c., exhibits no distinctive properties.

CHAPTER VI

THE FÆCES

UNDER the term "fæces"¹ are comprised all those substances which, being formed from the food in the process of digestion, and mixed with the residues of the secretions of the alimentary canal, are finally expelled from the body by the rectum.

I. NAKED-EYE CHARACTERS OF THE FÆCES.—The character of the fæces varies considerably in health, depending chiefly as it does on the nature of the food ingested. The labours of *Nothnagel*, however, have obtained for us a knowledge of some points which are more or less characteristic of healthy stools. Such a stool is moulded, and of a certain consistency. Its reaction is sometimes alkaline, sometimes acid. It is alkaline in certain morbid states, as, e.g., typhoid; but in others, as the acute enteritis of children (and, in the author's experience, of adults as well), the reaction is acid. The so-called "clayey" stools of dyspepsia in children, however, are nearly always strongly alkaline,—a fact which is due to the presence in them of carbonate of ammonia. *Nothnagel* concludes that there is little to be learned from the reaction of the fæces.

The colour, too, is very inconstant. It depends upon the food taken, and is greatly modified by drugs supplied to the system. When, e.g., bilberries are eaten freely, the fæces are coloured black. So, too, they are rendered black by preparations of iron, manganese, or bismuth, the colour in these cases being due to the formation of the sulphides of those metals. After a meal of cocoa-nibs or chocolate, the stools are apt to be coloured grey (*Widerhofer*).² The exhibition of calomel will turn them green, an appearance which was formerly attributed to the formation of sulphide of mercury, but which is now thought to be caused by the presence of biliverdin (*Betz, A. Vogel, Monti, Zaicadash*).³ Researches which the author has made with the green stools passed after the administration of calomel have shown that these contain abundance of urobilin, but no biliverdin. Hence it would appear that the colour is not due to the latter substance. *Lesage*⁴ distinguishes

two varieties of the green stool of children. The colour in the first is due to biliverdin, but in the second it is caused by a definite bacillus which produces a green colour substance. This bacillus can be cultivated outside the body, and inoculated upon animals. Its presence in great abundance in the fæces is associated with a severe form of cholera infantum. A green colour may also be produced by the *Bacillus pyocyanus* (*Koessel*,⁵ *Salus*).⁶ Santonin, rhubarb, and saffron will stain the fæces yellow.



FIG. 68.—Mucous Cylinder from the Fæces.

The presence of unaltered bile pigment in the fæces is always pathological (*Pettenkofer*).⁷ Healthy excrement contains a pigment which has been named stercobilin (*Vauvray*, *Masius*),⁸ and which *Maly* asserts to be hydrobile-rubin (urobilin). This latter body can be prepared artificially from bile pigment, and it is probable, *a priori*, that a similar change is effected in the process of intestinal digestion, when we would expect to find urobilin in the fæces. According, moreover, to investigations made by *G. Hopkins* and *A. Garrod*,⁹ Maly's hydrobile-rubin is not identical with urobilin, the former containing twice as much nitrogen as the latter. For a further description of this body and the tests by which it may be recognised, see the chapter on *Urine*.

The quantity of fæces passed by a healthy man in twenty four hours averages 120–200 grms. The remains of undigested food are often to be found in the excrement, as berries, fragments of potatoes and apples, and shreds of fibrous tissue. *Virchow*¹⁰ relates a case in which orange skins voided with the fæces were taken for a parasite, and *Eichhoerst*¹¹ one in which great coils of hard, woody asparagus were passed unaltered by digestion.

Amongst the occasional constituents of the fæces visible to the naked eye must be mentioned the cylindrical shreds of mucous membrane, of larger or smaller size, which are passed in the affection known as tubular intestinal catarrh (*Colica mucosa*, *Enteritis membranacea* or *tubulosa*), (*Nothnagel*).¹²

The cases in which such mucous formations are found are not very rare. Their discharge is attended with violent tenesmus, and unaccompanied by the passage of fæces. They are ribbon-like or reticular in form, and consist chemically of mucin and fibrin (*Litten*).¹³ They are

often of great length. In one instance they had the appearance of coiled brownish-yellow strings, 0.5 cm. thick, with here and there shreds of transparent membrane. The material under description was readily compressible beneath the cover-glass, and when looked at through the microscope exhibited a profusion of transformed intestinal epithelium involved amongst long, spirally-wound, and twisted threads. These bore some resemblance to the *Curschmann-Leyden* spirals (*vide* Chapter IV.), but had no central thread, nor did they exhibit crystals.

The specimen represented in fig. 68 was derived from a child of two years.¹⁴ It consisted of mucin and fibrin concentrically arranged, and enclosed bubbles of gas.

In all probability the presence of these bodies implies chronic catarrh of the large intestine, usually associated with obstipation and copious secretion of mucus.¹⁵

The author has observed similar appearances in the course of primary carcinoma of the pancreas. They have obviously no connection with the malignant growth.

In the case of a woman lately under his observation who suffered with peritonitis from perforation, the stools contained apparently similar bodies. These were seen microscopically to consist of fat and detritus. Chemical examination disclosed neither mucin nor fibrin. The substances in question had no resemblance chemically or microscopically to those mentioned above. The mucous lining of the gut was uninjured. It seems to follow that the condition known as *Enteritis mucosa* is in reality of a very variable character.

The author has seen in Vienna a body like in appearance to a tape-worm, and measuring $\frac{1}{4}$ metre in length, which was voided in a case which presently developed into chronic intestinal catarrh. Chemically it was found to consist mainly of fibrin and mucin. Further particulars as to the progress of this case are not to hand. *E. Henschen*¹⁶ describes a case wherein similar symptoms were caused by the presence of a species of fly akin to the common house-fly.

Virchow and *Nothnagel*¹⁷ describe the occurrence of bodies resembling frog-spawn or cooked sago-grains in the stools, and some observers have supposed that they were derived from ulcerating intestinal follicles. *Virchow* thinks they come from an excess of farinaceous foods. *Nothnagel* has also met with particles about the size of a poppy-seed, which chemically had the constitution of mucus. It must, however, be mentioned that this author has never found mucin in any quantity as a constituent of healthy excrement. *Kitagawa*¹⁸ found that these bodies in many cases consist of vegetable débris, but that others, which are distinguished by being more viscous and of softer consistency than the rest, are formed of mucus.

Foreign bodies of all kinds are occasionally to be seen in the fæces of lunatics and children.

Finally, tumours or parts of tumours originating in the alimentary canal, and stones or concretions formed in the gall-bladder or intestine, may be passed with the stools.

The presence of gall-stones is a fact of great clinical interest. They may in all cases be detected by a careful examination with the naked eye. They vary greatly in size, from that of a pin's-head to the size of a walnut. They vary also in consistence, being, however, less dense than the koprooliths which are occasionally met with.

For their detection it is necessary to wash the stools with water, strain and again wash. The friable substance which remains, may, but does not necessarily, consist of gall-stones. It may be formed of ingested substances, bony particles, gravel, &c., or of concretions (enteroliths) formed in the intestine. The latter have been described by A. Ott¹² and others.²⁰ Gall-stones are known by the determination chemically in them of cholesterin and lime. This is done by rubbing up a portion of the concretions in a mortar, boiling the powder in alcohol, and filtering. The filtrate is evaporated on a water-bath, and the residue submitted to the tests for cholesterin given in describing that substance in the present chapter. The part undissolved by alcohol is tested for carbonate of lime by the process described in Chapter VII.

The fæces of a woman suffering from various symptoms of dyspepsia were found to hold a number of soft, white and yellowish particles about the size of a pin's head. These proved to contain cholesterin and carbonate of lime, and it is probable that they were multiple small biliary concretions.

II. MICROSCOPICAL CHARACTERS OF THE FÆCES. To study the microscopical appearances of the fæces, a small particle should be pressed between a cover-glass and slide. If the stools are fluid, a drop may be placed upon a slide, and so examined. Much, of course, depends upon the character of the food, and the following description applies to the fæces of an adult living chiefly upon animal diet.

1. Constituents Derived from the Food.

(a.) **Vegetable Cells.**—These are of very variable form, and occur separately or in organic connection with one another. They sometimes exhibit starch granules or remnants of chlorophyll (fig. 69, *e-i, l*).

(b.) **Muscle Fibres** are almost invariably found in the fæces. Their quantity depends upon the supply of flesh-meat. They are relatively fewer after a mixed diet (*Nothnagel*).²¹ They are usually much altered, stained yellow by the bile, and swollen; but with a high power of the microscope, their transverse striation can usually be made out.

(c.) **Elastic Fibres** are readily to be distinguished by their double contour and curved form. They are always derived from the food, and occur both in health and disease.

(d.) **Connective Tissue** is found occasionally in the excrement of persons who eat much animal food, and is then of no pathological consequence. But if areolar fibres occur plentifully where flesh is sparingly eaten, they afford an indication of digestive disturbance.

(e.) **Fat.** Fatty globules are occasionally seen, but fat occurs more commonly in needles, arranged separately or in clusters (*Nothnagel*). Such needles are found in greater quantity when much fat has been taken with the food. The stools in alcoholic poisoning are always rich in fat, so also with the fatty diarrhoea of children (see p. 244).

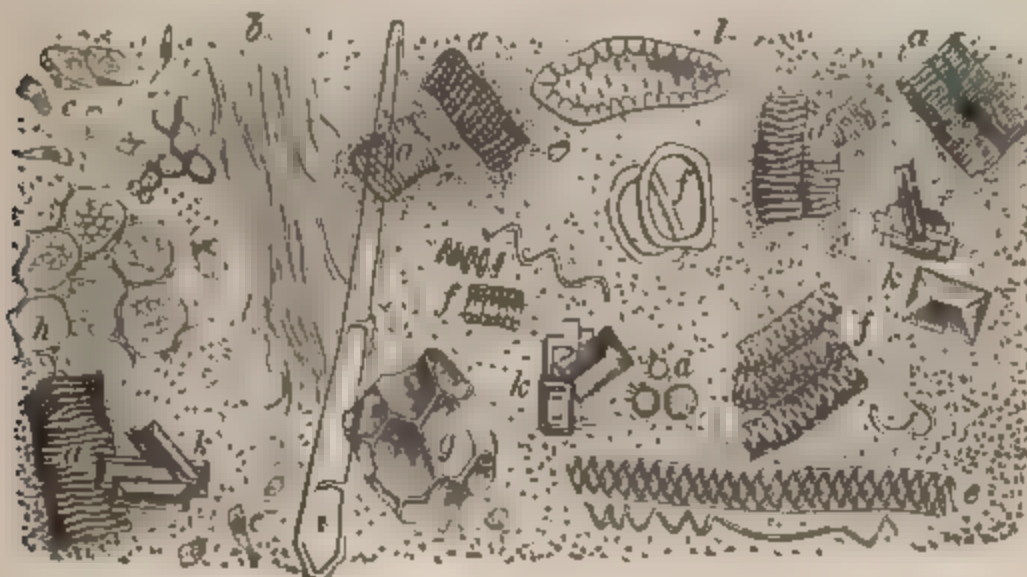


FIG. 69. Collective view of the Faeces (eye-piece III., objective 8A, *Reichert*).

a Muscle-fibres; b. Connective tissue; c. Epithelium; d. White blood-corpuscles; e. Spiral cells; f-i. Various vegetable cells; k. Triple phosphate crystals; l. Diatoms, the whole interspersed with a host of various micro-organisms.

(f.) **Starch Granules.**—These particles may be easily detected by their reaction with a solution of iodine and iodide of potassium. They are commonly to be found, but sparingly and of small size, in healthy stools. They are also found in the interior of vegetable cells. *Nothnagel* asserts that an excess of such substances in the excrement points to a derangement of the intestine.

(g.) **Coagulated Proteids.** Undigested milk occurs in the stools, especially in children and in persons suffering from diarrhoea. *Nothnagel* has described a substance resembling coagulated albumin which is occasionally to be seen in cases of intestinal trouble. It occurs in round yellow particles, varying from the size of a bean to that of a millet-seed. Their substance is readily soluble in a 5 per cent. solution of HCl, is precipitated from alcoholic solution by acetic acid, and rediss-

solves in excess of the acid, from which it may be again precipitated by ferrocyanide of potassium. They resemble the mucous particles of *Nothnagel*, to which reference has already been made. That author thinks that the substance in question is casein.²²

The discharges of infants at the breast are of a very different character from those of adults. Here muscle-fibres, areolar, and elastic tissue are absent, coagulated proteids taking their place. Microscopically they abound in fat and crystals of the fatty acid salts.

2. Formed Elements Derived from the Intestinal Tract.

1. Red Blood-Cells.—Red blood-corpuscles are rarely to be found in the fæces. *Nothnagel* has examined the freshly-voided excrement of typhoid, when it was deeply stained with blood, but could find none of these cells. In such haemorrhagic stools, however, are seen larger or smaller masses of reddish-brown pigment (haematoxin), and the rhombic



FIG. 70.—Degenerated Intestinal Epithelium (eye-piece II., objective 8A, *Reichert*).

crystals of haematoxin are also occasionally present. In cases where the blood is derived from the upper part of the alimentary canal or has remained for a long time in the intestine, the fæces are no longer red like blood, but are stained a dark-brown or black. We have already seen that a similar effect may be produced by certain drugs (see under "Naked-Eye Characters of the Fæces," above), and, consequently, it is not possible to infer a haemorrhage from this appearance alone; neither will red blood-cells be visible under the microscope. In such cases the presence of blood may be determined with absolute certainty by the application of *Teichmann's* test to a dried particle of the fæces (see Chapter I.).

2. Leucocytes.—In healthy stools leucocytes are rarely to be met with. They are always loaded with fatty particles. As a pathological condition, the appearance of leucocytes in great quantity in the excrement is far from common. *Nothnagel* found absolutely no increase in the number of these bodies in simple intestinal catarrh. When a

considerable increase does occur, it affords a presumption of ulceration of the gut.

Pure pus is found when an abscess has discharged into the intestine and in dysentery.

3. Epithelium.—Epithelium is always found in the faeces in health. Squamous epithelium comes from the region of the anus. Columnar epithelium is more rarely met with (fig. 69, c). The latter is usually uncoloured, but occasionally stained yellow. The cells occur either separately or in masses. Their boundaries, as a rule, are not easily to be recognised; but well-formed goblet-cells are sometimes seen (*Nothnagel*). Sometimes, too, they contain fat, and are then very large. Altered epithelial cells are often found in the faeces. In their typical manifestation they constitute what *Nothnagel* has termed “fusiform degeneration.” They are, for the most part, small, non-nucleated, homogeneous, and somewhat glossy bodies, tending towards the spindle-shape (fig. 69), but exhibiting every variety of form between this and the ordinary epithelial structure. *Nothnagel's* opinion is that these bodies are epithelial cells altered by abstraction of fluid, and he remarks that he has found them most typically shown in the mucous coating of scybalous masses. The mere presence of epithelium in the stools is a fact of no clinical consequence. When in disproportionate quantity, it points to intestinal catarrh.

4. Detritus.—There are certain indeterminate substances always to be found in the faeces, which are usually either derived from the food or belong to the waste products of digestion. They occur separately or in masses, and are little affected by reagents, although sometimes they are soluble in alcohol. Their constitution is very various.

3. Parasites.—There is no part of the body so apt to be infested by parasites as the intestine. Such parasites belong both to the animal and the vegetable kingdom, and it is not unreasonable to conclude, from the vast numbers in which some of them are always to be found, that they exercise an important function in the final processes of digestion. This may be said especially of certain vegetable forms, and among them the fission-fungi, presently to be mentioned, hold a prominent place.²³ [Researches by *MacFadyean*²⁴ point to a physiological distinction between the bacteria inhabiting the small and those of the large intestine. The former are stated to act only, or almost only, on carbohydrates, producing ethylic alcohol, which is constantly present in the small intestine; while the bacteria of the large intestine aid in the disintegration of proteids.]

A. Vegetable Parasites.—It will be useful to divide these parasites, as before, into two classes,—the pathogenic and the non-pathogenic; but in doing this, it must be premised that some of the organisms

classed as non-pathogenic may also at times be closely associated with morbid states. As an instance may be mentioned the *Bacterium coli commune*, which under certain circumstances acquires noxious properties (*Wyss*²⁵), giving rise not only to symptoms like those of typhoid, but to septic poisoning and to suppuration in various organs, as the kidney, liver, and bladder (see Chapter I.). The non-pathogenic parasites will first engage our attention, and we shall adopt the usual classification into moulds, yeasts, and fission-fungi.

1. Non-Pathogenic.

1. **Moulds.**—The only specimen of this class which has yet been found in the intestine is the thrush-fungus. It occurs in children suffering from thrush, and its presence in the stools is clinically of no significance.

2. **Yeasts.**—Yeast-cells (*Saccharomycetes*) are the commonest form of parasite in the intestinal discharges, whether of health or disease (*Nothnagel*), (fig. 69, between *b* and *c*). *Uffelmann*²⁶ asserts that yellow yeast-fungi are often also to be seen in the fresh stools of infants at the breast. Micro-organisms of this kind are most abundant in the acid stools of children. They are round or oval; lie together in groups of three or four; and commonly exhibit their characteristic sprouting arrangement. Well-formed yeast-fungi, however, such as are seen in fermenting saccharine solutions, are very rarely to be met with. *Nothnagel* saw such once in a case of typhoid in a child. In the bile-stained and acid discharges of acute catarrh of the small intestine in adults the author has not unfrequently found fungi which closely resembled the form described by *Rees*²⁷ as the *Saccharomyces ellipsoideus*, except that they were somewhat smaller.

The yeast-fungi of fæces stain a mahogany-brown with the iodo-potassic-iodide solution. This property depends upon the fact that they hold glycogen in their substance.

There are other forms to be met with morphologically resembling yeast-cells, but distinguished from them by giving a blue colour with the iodo-potassic-iodide solution mentioned below (see p. 201). The red and capsule-yeast, torula, &c., occur in meconium (*Escherich*²⁸), but are not known to have any morbid significance (see Chapter VIII.).

3. **Fission-Fungi.**—These organisms exist in swarms in the intestine, and they are to be found in greater profusion in the fæces than in any other of the excretions (*Nothnagel*, *Brieger*, *Uffelmann*, *Escherich*, *Bienstock*, *Stahl*, *Kuisl*, *Miller*, *Sucksdorf*²⁹); indeed, it is not incorrect to say that they always constitute the bulk of these discharges. Bacilli and micrococci of the most varied kinds are the commonest forms. They occur separately or in colonies, and often exhibit lively movements. It may be said, as a rule, that micrococci predominate in the solid, and

bacilli in the fluid motions. The former are sometimes found in shapes resembling torulæ and sarcinæ. The organism which occurs most commonly and in greatest abundance is the *Bacterium coli commune*, and there can be little doubt that this body is intimately associated with putrefaction processes in the intestine; although it must be stated at the same time that we are ignorant of its functions in health, where also it exists abundantly, and that the host of other micro-organisms which make their appearance when putrefaction is going on throws a great deal of doubt over this question. It must once more be mentioned that this bacillus as an infective agent is at times among the most formidable of human parasites.

Bacillus subtilis is found in the discharges both of health and disease. It was first discovered by *Nothnagel*, and may be seen as long threads with spores attached, or as separate spore-bearing rods, and elsewhere the spores alone occur in clusters. Its relatively thick edges and the remarkably glossy appearance of the spores facilitate its detection. Its presence has no clinical significance.

The various micro-organisms above alluded to stain brown or brownish yellow in solution of iodine and iodide of potassium or of ammonium iodo-iodide; and this property belongs especially to the groups of micrococci, which are coloured very deeply a brownish yellow by contact with this reagent.

In addition to such forms, the fæces exhibit other micro-organisms, which stain blue or violet in the iodo-potassic-iodide solution. *Nothnagel* has described many of these, and one especially which he holds to be identical with the *Clostridium butyricum* of *Prazmoicshi*.³⁰

The author has investigated this subject with great care, and he is in a position to confirm *Nothnagel's* statements in every particular, and would venture to add something to his description of the fungi which stain blue with the iodo-potassic-iodide solution.

To begin with the minutest forms. There is a micrococcus which occurs in swarms of uniformly and finely granular zooglœaform bodies, which colour a reddish violet with the solution.

Next in order is a micro-organism in the form of short delicate and somewhat pointed rods, which recall the *barillus of septicæmia in the mouse*, and which stain in a similar manner with the reagent mentioned. These rods occasionally contain one or two spherical granules, which do not stain in the solution.

There are other rods, of varying length, which resemble the *Leptothrix buccalis* in the manner of their reaction with the iodine fluid; and a further variety, which differs from the *Bacillus subtilis* only in that the fungus threads stain blue, while the bodies referred to above as the spores remain uncoloured (fig. 71).

The author has often had an opportunity of observing the *Clostridium butyricum* of *Nothnagel*. It takes the form of large round cells, usually visible in uncoloured preparations by reason of their somewhat lustrous appearance, and in other respects resembling yeast-fungi. In some cases they adhered to one another like a string of beads, and in others were simply disposed in groups (fig. 72). These bodies, as *Lichtheim* and



FIG. 71.—Bacilli from the Fæces, staining Blue with the Iodo-Potassic-Iodide Solution (eye-piece III., objective 8A, Reichert).

others have observed, stain with the Ziehl-Neelsen fluid in the same manner as tubercle bacilli; but they are not likely to be mistaken for these, since they are sufficiently distinguished by their size, shape, and peculiarity of arrangement.

A micro-organism of oblong and somewhat pointed form also occurs in the fæces, and cultivations of this have been made by *H. Fischer*.

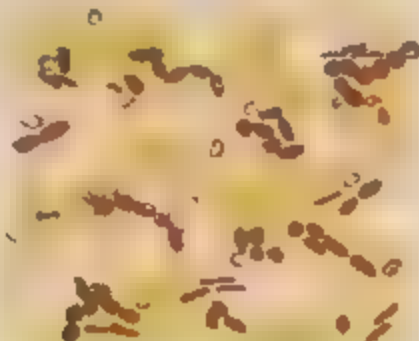


FIG. 72.—*Nothnagel's Clostridia* and Stunted Bacilli from the Fæces, staining Blue with the Iodo-Potassic-Iodide Solution (eye-piece III., objective 8A, Reichert).

It must be mentioned that although these bodies all stain some shade of blue with the solution of iodine and iodide of potassium, the tints which they derive from that reagent exhibit notable differences, for whilst, on the one hand, the micrococci are but slightly coloured, and of a purple tint which tends towards red, the rod-like micro-organisms stain very deeply, and of a dark-blue colour. The stain is in all cases transitory, fading in from twenty-four to forty-eight hours, and disappearing altogether within a few days.

Those fungi whose characteristic it is to stain blue in iodo-potassic-iodide solution occur probably in all stools, but in relatively small numbers in those of health. They are in profusion only in certain morbid states, and especially in intestinal catarrh. They do not seem to bear any relation to the reaction of the discharges, having been found alike when this was alkaline and when it was acid. To conclude, they are constantly present in health and in disease, and in the discharges of infants at the breast, as well as of older children nourished on a meat-diet. By Koch's processes many, though not all, of the micro-organisms occurring in the fæces can be isolated and studied. It is certain that some of them are pathogenic.³¹

Hence it follows that no definite clinical significance attaches to any of the parasitic forms hitherto described, with the exception of the *Bacterium coli commune*. It is a matter of observation that in certain pathological conditions of intestinal derangement one form or the other is apt to preponderate, but we have no evidence that they are in any case the cause of disease, or that the multiplication of a particular micro-organism is not rather the consequence of the disturbance which it is found to accompany.

The intestine is apt to be infested with microbes of a pathogenic character, which in form closely resemble those innocuous parasites which have just been described. Much light has been thrown upon their nature by the researches of recent years, and for their detection we have to make use of a number of special methods. But these alone will not suffice. Without an accurate knowledge of the commoner, at least, of the non-pathogenic organisms which normally inhabit the intestine, the discrimination of the others is impossible. It is for this reason that we have dwelt at some length—though still in a far from exhaustive manner—on the description of the innocuous micro-organisms which are most frequently to be found in the fæces.

2. Pathogenic Fungi.—We pass now to a consideration of the pathogenic parasites, the bacilli of cholera, typhoid, and tubercle.

1. Cholera-Bacillus (*Comma-Bacillus*).—To *Robert Koch*,³² the pioneer of the modern science of bacteriology, belongs the honour of having first discovered the micro-organism which causes that most terrible of the epidemic diseases of our time—cholera.

We shall not attempt to give an account of all that has been written upon this subject, but shall content ourselves with making here and there a reference to the leading authorities for the information of the reader; and we shall avoid the discussion of controverted points of which we have no personal knowledge. *There can be no doubt whatever of the fact that a definite and morphologically distinctive parasite occurs in the discharges of cholera patients.* Nevertheless, the knowledge

recently acquired²³ in well-investigated cholera epidemics has conclusively shown that this micro-organism is sometimes absent in typical cases of the severest form of cholera, that at such times the cholera-bacillus is harboured in the intestines even of healthy individuals, and that, more over, a whole series of micro-organisms (such as *Vibrio danubicus*) exist which have a close morphological relationship to the cholera-bacillus, and that it is difficult to distinguish these by the usual methods of cultivation. These facts do not lessen the significance of the cholera-bacillus, but they limit and complicate our diagnostic resources.²⁴ Nevertheless, it is the duty of every physician to be confident as to their indications.

Koch described the cholera-bacillus as a short rod-like form, curved or semicircular, and somewhat thicker than the bacillus of tubercle. Such bodies often lie two together, with the concavity of their curves turned in opposite directions and their extremities in contact, so as to form an S-shaped figure. *Neuhauß*²⁵ describes flagella upon them.

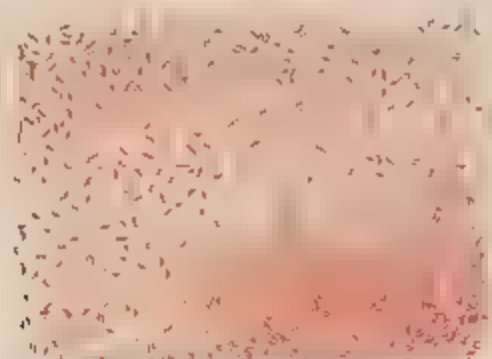


FIG. 73.—Koch's Comma-Bacillus (pure cultivation; eye-piece III., objective Zeiss 3, homogeneous immersion, Abbe's mirror and open condenser).

*Löffler*²⁶ has demonstrated their existence by the method described in Chapter X. They produce by division peculiar screw-like spirals, which remind one of the spirillum of relapsing fever, only that these are thicker (fig. 24). Koch did not find any separate spores belonging to these organisms, but it would seem that *Hueppe*²⁷ has succeeded in doing so. Such bacilli were discovered by Koch in the intestine and discharges, rarely in the vomit, of Asiatic cholera, and never in the blood, saliva, tears, urine, or breath of patients suffering from this disease. The discharges, according to this authority, sometimes contain almost unmixed colonies of bacilli. Koch's statements have been borne out by the subsequent investigations of *Baber*, *Vandyke-Carter*, *Nicati*, *Rietsch*, *van Ermengen*, and others.²⁸

It follows from what has been said of the enormous profusion in which micro-organisms of various kinds infest the intestine, that the cholera-bacillus, when not very plentiful, may easily escape detection, and therefore it will not suffice to submit discharges which are supposed

to contain it to a simple microscopical examination. To meet the exigencies of the case, Koch has studied the mode of growth of the bacillus, and he has devised for its cultivation certain special processes, on the application of which the diagnosis of Asiatic cholera may very safely rest. The first requisite is to separate some of the bacilli or their germs from the mass which contains them, and this may readily be done in the manner to be described in another part of this volume (*vide Chapter on Methods of Bacteriological Research*).

A complete and exhaustive search for the cholera-bacillus may be performed as follows :—

1. A particle of the discharge is placed upon a slide, and examined microscopically for the bacillus. In doing this, the expedient of *Schottelius*³⁹ may be adopted with advantage. He puts a little of the stool in an open glass, together with an equal quantity of alkaline meat-broth, and allows the mixture to stand at a temperature of 30°-40° C. for twelve hours. At the expiration of that period the bacilli will be found in swarms on the surface of the glass, and a specimen may be obtained which will consist almost entirely of them.

2. A particle of the discharges or a drop of the infected broth (*Schottelius*) is next spread out in a very thin layer between two cover-glasses, dried, passed three times through the flame of a Bunsen burner, stained with one of the basic aniline dyes (fuchsin, methylene blue), and examined under the microscope. According to Koch,⁴⁰ a simple inspection of a cover-glass preparation directly from the stools stained with carbol-fuchsin (see p. 128) suffices to establish the diagnosis. In such a preparation the micro-organisms are seen massed together like a shoal of fish in still water. This appearance is said to be characteristic.

3. Plate-cultivations are made from the suspected stools with nutrient gelatine and agar-agar, as described in Chapter X.

4. If comma-bacilli develop in these, cultivations are to be made by inoculation in the depth of nutrient substances.

5. The bacillus is to be cultivated in hanging drops (*vide Chapter X.*—it is well to apply this test at once, in the event of comma-bacilli being found in Schottelius' preparation)—and the micro-organisms developed in the medium are to be compared with those derived by plate-cultivation.

6. Inoculation is made on animals with the pure cultivation.

7. The peptone cultivation is tested for the indol-reaction.

When the discharges examined belong really to Asiatic cholera, the processes 1 and 2 will generally show the comma-bacillus of Koch. The inference to be drawn from the third process depends upon the fact that the cholera-bacillus, cultivated on a nutrient gelatine plate at

22° C., forms, after twenty-four hours, white colonies, with irregular, jagged, or sinuous outlines. These exhibit a light-yellow or rosy tint, and present an appearance like that of a layer of powdered glass overlying the gelatine plate. The colonies grow gradually darker towards their centre, and presently begin to liquefy.

Plate-cultivations in nutrient agar-agar form a greyish-yellow, furrowed, slimy surface, and do not cause the underlying nutrient medium to liquefy.

When cultivated by inoculation in the depth of the nutrient substance contained in a test-tube (*v. supra*), after the lapse of twenty-four hours the nature of the bacillus is shown by a white coloration, which extends along the track of the needle, and the formation of a funnel-shaped cavity, which increases gradually from the circumference, and acquires the appearance of an air-bubble. It is only at its superficial part that this liquefaction occurs, the lower layer of the inoculated substance remaining unaltered for days.

In hanging-drop cultivations, the comma-bacillus exhibits the following peculiarities (*v. supra*) :—When examined on the day following the inoculation, or, it may be, after the lapse of a few hours, and with an oil-immersion lens and a constricted diaphragm, mobile swarms are seen in the centre of the mass, while at its circumference appear the spirochæte-like bodies, which sometimes exhibit as many as twenty spiral twists. Suppose that a specimen, which Schottelius' method of Koch's peptone cultivation had previously shown to contain micro-organisms when cultivated in this manner, displayed some forms which resembled the comma-bacillus, it will then be necessary to transfer some of these drop-cultivations in the manner described in Chapter X. for cultivation by the plate and deep-inoculation methods.

Bujicid has recently recommended his chemical process (see below), in combination with that of Schottelius, for the purpose of disclosing the cholera microbes, even without the aid of the microscope.

Further confirmation may be had by the experiment on animals (No. 6). A particle (1.5 mgrm.) of the cultivation from agar is taken on a platinum point, suspended in 1 cc. of bouillon, and injected into the peritoneal cavity of a guinea-pig, where it induces a typical toxic effect. Koch⁴⁰ attaches much importance to this process, because, of all the spirilla-like bacteria, that of cholera is the only one which induces the symptoms in question.

It should be mentioned that the cholera-bacillus will thrive at a temperature of 37°, and even on boiled potatoes. Its cultivations resemble those of the bacillus of glanders to the naked eye, but their growth is slower and needs arti-

ficial heat. The bacilli are very sensitive to drying and to exposure to a 5 per cent. solution of carbolic acid.

Bitter and *Rietsch*⁴¹ have shown that the cholera microbes elaborate a peptonising ferment.

Poehl and *Bujwid*⁴² found that the addition of a 5 to 10 per cent. solution of hydrochloric acid will, in a few minutes, impart to cholera cultivations, and to no others, a violet-rose colour; and *Brieger*⁴³ has been able to separate from them a colour-substance, to which he gives the name of "cholera-red," but which *Salkowski*⁴⁴ identifies with indol. *Bujwid's* cholera reaction does not seem to merit the commendations bestowed on it. Other fungi, pathogenic and innocuous, behave in a similar manner with mineral acids.

The foregoing observations (which were described by the author as long ago as in the second German edition of this work) were confirmed by *Kitasato's*⁴⁵ earlier work; but since they were first made, the whole matter has acquired a more solid basis from *Koch's*⁴⁶ admirable researches, which have shown that the colour reaction is not produced by any other of the curved bacilli but that of cholera.

To obtain it *Koch* proceeds as follows:—A cover-glass preparation is first made, and the bacilli being identified, a cultivation is made on peptone at 37° C., from which a pure cultivation is obtained after the lapse of eight hours. The addition of a nitrate and pure sulphuric acid should develop the indol reaction. A pure cultivation only should be employed, the presence of a nitrate in the peptone is indispensable, and the acid must be free from sulphurous acid.

The pure cultivation derived from peptone may now be transferred to the gelatine plate, and in twenty hours at 22° C. the appearances described at p. 206 may be noted. Another cultivation may be made on agar-agar, and after eight to ten hours at 37° C. examined with the microscope (*vide supra*); and finally, inoculation upon an animal may be resorted to.

In view of the immense importance of recognising the nature of an epidemic of cholera at the outset, every officer of the public health should make himself acquainted with these processes, which undoubtedly are invested with the largest measure of certainty.

Cantani notices, as a result of his experiments upon animals, that cholera-bacilli elaborate a poison; and *Brieger*⁴⁷ has actually separated from its cultivations specific toxic substances, which occur together with cadaverin and putrescin, and result from the agency of cholera microbes. The method he employed is that detailed at p. 187. It remains for the clinical observer to detect these substances ready-formed in the discharges of cholera patients; and some progress has already been made in this direction (*Pouchet, E. Roos*).⁴⁸ Recent research has made it doubt-

ful whether it is these toxines which play so important a part in the morbid process, and not rather the toxalbumins from which they are known to originate.

There are other micro-organisms which bear a considerable resemblance to the comma bacillus. One of these, the bacillus of cholera nostras, is a pathogenic micro-organism; another is Deneke's spirillum of cheese, and a third is the Vibrio danubicus of Heider.

2. The Bacillus of Cholera nostras.—*Finkler* and *Prior*⁴⁹ have observed a micro-organism resembling the comma-bacillus in the discharges of cholera nostras. It is distinguished from the bacillus of Asiatic cholera chiefly by its size (fig. 74., being both longer and broader than the latter. In addition to this the two exhibit notable differences in life-history. When a colony of the *Finkler-Prior* bacillus is cultivated on a plate in nutrient gelatine, it is seen to be of a uniformly round figure with well defined edges; and when examined with a low or medium power, it has a granular appearance, and is usually of a brown colour. Moreover, the gelatine rapidly liquefies, and emits a very foul and penetrating odour.



FIG. 74.—*Finkler-Prior* Bacillus of Cholera nostras (pure cultivation; eye-piece III. objective Zeiss $\frac{1}{2}$, homogeneous immersion, Abbe's mirror, open condenser).

The comma-bacillus of *Koch*, on the other hand, develops less rapidly. The colonies are never coloured brown, but are of a tint ranging from light yellow to rose-colour; and the figure which they form, as already mentioned, is bounded with a ragged contour (see p. 206).

Again, when cultivated by inoculation in the depth of nutrient substances, this bacillus develops in a very characteristic way; for whilst the comma-bacillus swarms in the funnel-shaped manner already described, the cultivation in this case assumes a saccular shape, somewhat comparable to that of a stocking. *r. Horovka* and *Winkler*⁵⁰ employ plover's egg-albumin as the food medium. This is rapidly liquefied by the *Finkler-Prior* bacillus, while the comma-bacillus is propagated only along the track of the needle and does not decompose it.

The pathological import of the *Finkler-Prior* bacillus is still a matter of dispute, but it is plainly of the utmost consequence to be able to distinguish this comparatively harmless organism from the very fatal microbe of Asiatic cholera.

Spirillum of Cheese.—*Deneke*⁵¹ found a micro-organism in ripe cheese which bears a close resemblance to the comma bacillus of *Koch*. But this, like the *Finkler-Prior* bacillus, can be distinguished by certain peculiarities in its life-history. The nutrient gelatine medium is rendered fluid sooner than by *Koch's* bacillus, but more slowly than by that of *Finkler* and *Prior*. This micro-organism, moreover, will not develop in potato, while the other two will thrive

upon this food. The crucial test, however, is the result of inoculation upon animals. Deneke's bacillus has no morbid influence on the intestine.

Dumbar,⁵² Dergel,⁵³ and Rumpel,⁵⁴ who examined the water of the Elbe at the time when the cholera epidemic was raging in Hamburg, discovered a bacillus whose characters closely resembled that of cholera, being distinguished from it only by a more rapid development in nutrient media. Ralner⁵⁵ reported similarly.

Hader⁵⁶ discovered in the water of the Donau Vienna Canal, at a time when cholera was not prevalent, a bacillus very closely resembling that of cholera. He named it the *Vibrio danubicus*. It was probably the same described by C. Frankel.⁵⁷ Its import is not yet known, but it seems to be either a variety of the cholera-bacillus or a closely related organism.

3. Bacillus of Typhoid Fever.—A characteristic parasite was discovered in 1880 by Eberth⁵⁸ in the implicated viscera in typhoid.



FIG. 75. Bacilli of Typhoid (pure cultivation—eye-piece III—homogeneous immersion, objective Zeiss 1½). From a preparation by Dr. Paltau.

Similar observations were afterwards made by Klebs and Eppinger,⁵⁹ and these have been confirmed by the researches of R. Koch, Meyer, and Friedlander, and more recently by Gaffky and very many others.⁶⁰

Gaffky describes the fungus as consisting of rods, in length equal to one third the diameter of a red-blood corpuscle, and occasionally forming threads of greater length by the aggregation of several segments. They are about three times as long as they are broad, and rounded off at the extremities. Spores are sometimes to be seen within the rods. They stain best in a concentrated watery solution of methylene blue, and Loeffler's process (v. Chapter I.) is the most appropriate to the purpose. They are not stained by Gram's method. Frankel and Pfeiffer,⁶¹ using Leijlet's process for staining flagella, have observed these processes on the bacilli of typhoid. Similar flagella have been observed upon other micro-organisms, and notably on the *Bacterium coli communis*.

Gaffky⁶² has ascertained certain facts connected with the life history of the typhoid-bacillus. It develops readily in peptone-gelatine bouillon.

In a preparation of this kind the cultivations make their appearance after the lapse of twenty-four hours. When examined with a medium power of the microscope, they seem to be yellowish, and they do not render the gelatine fluid. Rods and threads appear, and they are endowed with an evident and peculiar motion. The parasite develops on prepared potato. The colonies can be distinguished with difficulty by the naked eye. When cultivated in a medium of prepared potato at a temperature of 37° C., spores begin to form after three or four days. According to *Birch-Hirschfeld*,⁶³ solitary and segmented spores are to be seen, the first in hanging-drop cultivations, and the latter when cultivated in an incubator. This observer recommends that the nutrient medium be stained with phloxin-red or benzo-purpurin, by means of which the spores are deeply coloured. There is much doubt as to the true character of these so-called spores (*Buchner, Pfeuhl*).⁶⁴

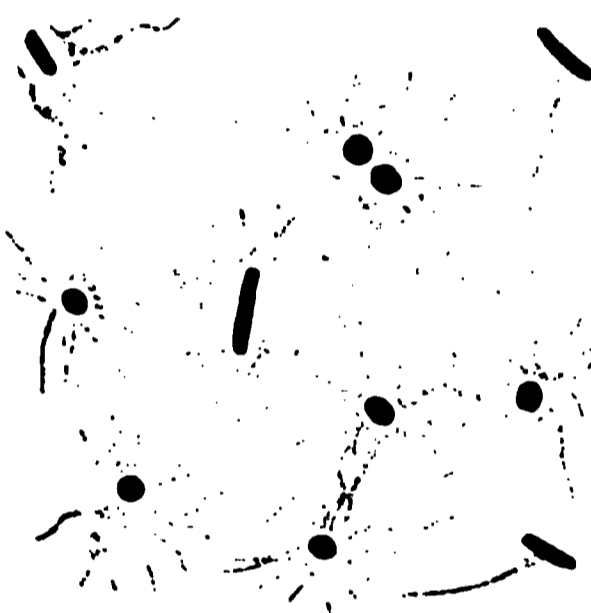


FIG. 76.—Bacilli of Typhoid with flagella. (Zeiss's compensation objective VIII., apochromatic immersion 1.30). From a preparation by Chiari.

The fungi are easily cultivated in hanging drops in sterilised broth. They develop in nutrient substances containing carbohydrates, as cane or grape sugar or lactose. They fail to give the indol reaction in albuminous media.⁶⁵

The bacillus occurs in the stools of typhoid patients; but on account of the vast number of micro-organisms constantly to be found in the dejections, it is impossible to recognise it with the aid of the microscope alone. Neither does it possess, like the bacillus of tubercle, any characteristic staining properties. For its complete recognition, therefore, we must resort to Koch's methods for obtaining pure cultivations, and separate it from the faecal substances. *Pfeiffer*⁶⁶ was the first to do this, and he made use of the nutrient agar-agar plate. A useful adjunct is the employment of a 0.25 per cent. solution of carbol-gelatine, after the manner of *Chantemesse* and *Widal*.⁶⁷ According to *Holz*,⁶⁸ however, this is impracticable, since he found that the development of

the bacilli was suspended in presence of 0.1 per cent. of carbolic acid. The best medium is neutral potato-gelatine containing 0.05 per cent. of carbolic acid. For the recognition of this micro-organism, *Holz*, following *Grancher* and *Deschamps*,⁶⁹ recommends weakly-acid bouillon or similarly-prepared milk stained by *Noeggerath's*⁷⁰ process. *Kitasato*⁷¹ relies upon the absence of the indol reaction from such a cultivation as evidence of the typhoid-bacillus.⁷² It is especially difficult to distinguish the typhoid-bacillus from *Bacterium coli commune*. *Lyonnet's*⁷³ process for doing this is as follows:—Ordinary bouillon (see Chap. X.) is stained with animal charcoal, and to it are added 1 per cent. carbolic acid, 2 per cent. sugar of milk, and a little Congo-red. Upon this is implanted the suspected material from the stools. Upon this medium there develop only the bacillus and the *Bacterium coli*. If the medium remains transparent, neither were present. If it becomes turbid, but remains red, then probably there is only the bacillus there. Should it become turbid, and also turn violet (from the presence of free lactic acid), the *Bacterium coli* is probably involved. *Marpmann*⁷⁴ proposes to discriminate between these micro-organisms by adding reduced colour-substances to the nutrient media. *Elsner*⁷⁵ advocates the following method:—Ordinary gelatine is boiled with potato extract ($\frac{1}{2}$ kgrm. to a litre of water); 2.4 to 3.2 cc. of a normal caustic soda solution (*Holz*)⁷⁶ added to secure the necessary point of acidity, and the fluid is filtered and sterilised. For the cultivation of typhoid bacilli the gelatine is poured into an *Erlenmayer's* flask, 1 per cent. of potassium iodide added, the mixture inoculated, and the requisite plates run off. After twenty-four hours colonies of the *Bacterium coli* will have developed, while forty-eight hours must elapse before the small shiny water-drop-like bodies show the presence of typhoid cultivations. *Brieger*⁷⁷ very highly commends this process for clinical purposes. *Pollak*⁷⁸ tested the method in the author's clinic, and found it possible by this means to speedily separate *Bacterium coli*, *Bacillus typhi*, and *Bacillus faecalis alcaligenes*. He considers the method useful, but, so far as the author's experience extends, it does not fulfil the claim put forward by *Brieger*, viz., the rapid and reliable detection of the typhoid-bacillus. The researches of *Babes* and *Cassedebat*⁷⁹ have shown that the investigation is attended with much difficulty, since there are a great many distinct fungi which, when cultivated, present appearances quite similar to those of the typhoid-bacillus.

It is difficult to effect the separate cultivation of typhoid-bacillus derived from the faeces in nutrient gelatine, because the discharges contain other micro-organisms (hay-bacillus), which liquefy that medium before the colonies of typhoid-bacilli appear in it.

When the micro-organism under discussion has been experimentally communicated to animals, it causes them to manifest the symptoms of

typhoid; and the researches of *E. Frankel, M. Simmonds, and C. Seltz*,⁸⁰ seem to leave no doubt as to its pathogenic character. *Beumer and Peiper*,⁸¹ on the other hand, have come to a different conclusion. Bearing in mind *Brieger's* discovery of animal alkaloids (*ptomarines*) as a product of bacillary life, it seems probable that such play an important part in connection with typhoid, and this may well explain the controverted results obtained in the experiments upon animals reported by *Frankel, Simmonds, and others*. [The toxines obtained from the urine of typhoid patients are said to resemble very closely in their characters those isolated from artificial cultivations (*Luff*).]

[**The Sero-Diagnosis of Typhoid.**—1. *Historical Survey*.—The experiments of *Widal*⁸² have secured a most important aid to the rapid diagnosis of typhoid. His method is based upon the peculiar properties possessed by the serum of immunised animals, properties which may be generally described as toxic to or inimical to the growth and development of the specific micro-organisms concerned. The bactericidal power of such serum has been established by numerous test tube experiments, as those of *Charren* and *Royer*⁸³ in the case of the *Bacillus pyocyanus*, of *Behring and Nissen*⁸⁴ with the *Vibrio Metchnikovii*, of *Metchnikoff*⁸⁵ with the pneumococcus. *Pfeiffer*⁸⁶ found that the injection of cultivations of the *Cholera* vibrio into the peritoneal cavity of immunised guinea-pigs was followed by degeneration or death of the micro organisms, and subsequently, working with the *Bacterium coli* and the bacillus of typhoid, obtained a similar result. *Bordet*⁸⁷ extended and improved upon *Pfeiffer's* method. He added a little of the protective serum to debrinated blood from healthy guinea-pigs, and observed that the mixture when fresh has the bactericidal property; and, even when preserved for a time, he found that, while not destructive of the micro-organisms to which it was fatal when fresh, the fluid had a peculiar effect upon these, stopping their motion and causing them to come together in masses or clumps. *Gruber*⁸⁸ and *Durham*⁸⁹ further studied this phenomenon, and defined the conditions in which it occurred most typically. While *Bordet* showed that normal serum of certain animals (as that of the horse) could cause the agglomeration of particular specific micro-organisms, *Durham* ascertained that the "sera which react most strongly are those which are prepared by virulent cultures of the same species," and that the reaction is not given by the *Bacterium coli* with the serum of animals immunised against typhoid fever, and vice versa, and he used this method to differentiate between these micro-organisms. The principle of the specificity of the "clumping" phenomenon would seem to be established by these facts.

Chantemesse and *Widal*,⁹⁰ and afterwards *Brieger, Kitamoto, and Wassermann*,⁹¹ have investigated the experimental immunisation of animals against typhoid fever. *Stern*⁹² employed the serum of convalescent typhoid patients in similar experiments, and, finally, *Chantemesse* and *Widal*⁹³ ascertained that the serum of persons who had had typhoid had a curative and protective action in respect of the disease produced experimentally in animals.

2. *Widal's Method.* *Widal*⁹⁴ discovered that the blood serum of typhoid patients

* For the statements in the text and the references the editor is indebted to an admirable paper by *S. Delepine* in the *Med. Chronicle*, Oct. 1896, New Series, vol. vi.

'Author's Note.—Out of regard for my deceased friend *Cagney*, this short comment on Sero-Diagnosis is retained in this edition, although the subject has been dealt with by the editor in Chapter I.]

has the same action upon cultures of the typhoid-bacillus as has the serum of animals immunised against typhoid; and further, that the serum in this case has the property of causing the "clumping" phenomenon, even when taken at a very early period of the disease. He devised a method, at once simple and secure, by which this property was made the basis of an admirable clinical test. It is applied as follows:—(1.) The patient's finger (thoroughly cleansed and sterilised) is pricked, the blood received in a sterilised glass cell, and the serum allowed to separate. (2.) One drop of serum is placed on a sterilised glass slide. (3.) To this is added ten drops of a recent pure cultivation of typhoid-bacillus. (4.) The mixture is covered with a cover-slip and inspected under the microscope at once. If the reaction is not evident, the specimen is examined again at intervals up to two hours.

The reaction consists in this, that the bacilli run together and gradually become motionless. The "clumping" and loss of motion, although they occur together, may in the present state of our knowledge be profitably distinguished; and some observers attach more importance to the latter than to the former.

*Delépine*⁹⁵ has employed this method in a large number of cases, both of typhoid and of other diseases, with complete success. He proceeded thus:—(1.) The finger (cleansed and sterilised) is pricked, and a little of the issuing blood is sucked into a modified *Pasteur's* pipette, previously sterilised. The point of this is then sealed, and the constriction of the pipette is drawn out in the blow-pipe and also sealed. When the blood is about to be used, the point of the pipette is passed through a flame and broken off, and the serum (expelled by heating the broader end) is received on a sterilised slide. (2.) Part of a pure cultivation of typhoid-bacillus (24–48 hours old) in neutral peptone bouillon is poured into a sterilised watch-glass, and from this, with a platinum loop 1 mm. diameter, nine drops are taken, and placed separately, but near each other, on a cover-glass. (3.) The loop is again passed through the flame, and with it one drop of the serum is placed on the cover-glass, and the ten drops are mixed and inverted on a slide. (4.) The preparation is examined after the lapse of one, two, or three minutes, and, if necessary, at intervals of an hour up to twenty-four hours. After the first few minutes it should be kept in a moist chamber.

Widal has obtained the reaction with serum taken as early as the fourth day of the fever, and after the eighth week. *Delépine* secured it most easily between the end of the first and the end of the fifth week. In twenty-five cases of typhoid it never failed, and in ten cases of disease which were not typhoid it never appeared (*Delépine*).⁹⁶ *Chantemesse*⁹⁷ has reported similarly. *Widal* and *Sicard*⁹⁸ found that serum was more effective than blood. Dry serum will serve, but loses its power more quickly. *Achard* has seen clumping under the influence of milk from a woman affected with typhoid, and *Widal* records varying results from the use of typhoid urine.]

Finally, it may be noticed that many recent investigations tend to prove the possibility of typhoid-bacilli being disseminated by means of drinking-water and by milk.⁹⁹

4. Bacillus of Tubercle.—*Lichtheim*¹⁰⁰ and other observers have found tubercle-bacillus in the stools in cases of tubercular ulceration of the intestine. For its recognition there, the methods are employed which have already been described in connection with the sputum, *vide Chapter IV.*

The detection of this bacillus in the faeces invariably implies the existence of tubercle, but not necessarily tubercle of the intestines, since

it may be derived from sputa which have been swallowed. Still, in cases where it has been repeatedly found in the stools, and where it occurs in great profusion, so as to resemble pure cultivations (*vide* fig. 60), and, especially, where the other symptoms of tubercular ulceration of the intestine (purulent discharges, &c.) are present, the diagnosis is established on the firmest basis.

5. *Bacterium coli commune*.—Much interest and importance has lately come to be attached to the presence in the alimentary canal of this micro-organism, and it has been seen (*vide* Chapter I.) to be a matter of consequence to be able to distinguish it from the typhoid-bacillus, with which it is readily confused.

For its detection *Koch's* method may be adopted. The cover-glass preparation shows rods of varying length, which stain readily in dilute carbol-fuchsin fluid. By a staining appropriate to the purpose (*vide* Chapter X.) the bacterium may be seen to have as many as three

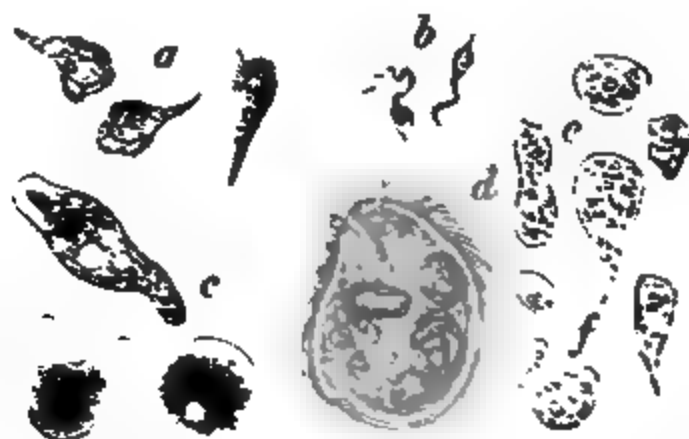


FIG. 60.

- | | |
|---|-----------------------------|
| a. <i>Trichomonas intestinalis</i> . | d. <i>Paramecium coli</i> . |
| b. <i>Cercomonas intestinalis</i> (Dien.) | e. Monadines, living. |
| c. <i>Amaroba coli</i> . | f. Monadines, dead. |

flagella. It stains by *Giemsa's* method only, when it has developed in a fatty medium. On nutrient agar the cultivations form a whitish coating, and on the gelatine plate they penetrate deeply and have a radial or concentric arrangement. The gelatine is not liquefied. On potato chips they form a thick, shining, brownish layer. They curdle milk by the evolution of lactic acid. The micro-organism is polymorphous, and many authorities are of opinion that several distinct organisms are here confused under one name.

B. ANIMAL PARASITES.

1. Protozoa.—Adopting the classification of Leuckart,^{1,2} these include Rhizopoda, Sarcodina, and Infusoria.

1. Rhizopoda.

(a) **Monadines.**—Nicolson³ found these organisms repeatedly in

the stools of consumptive and typhoid patients, and of persons suffering from heart-disease. They were always dead unless the stools were examined directly upon being passed, and then appeared as more or less circular bodies of various sizes (fig. 77, *f*).

The still living monadines are pear-shaped, and usually possess a long pointed process, which moves about rapidly (fig. 77, *e*). According to *Nothnagel*, they have no pathological significance. *Grassi*¹⁰⁸ found similar bodies in the stools of a patient suffering from entero-colitis. Bodies altogether resembling these, but not as yet identified with them, occur in the discharges of infants and children (*v. Jaksch*).¹⁰⁴

(b.) *Amœba coli*.—*Lösch*¹⁰⁵ has described certain cellular bodies of large size which he found in the faeces in a case of intestinal ulcer of a quasi-tubercular character. These bodies were contractile, and some of them, of circular form, had a diameter of 20–35 μ . They consisted of hyaline and coarsely granular protoplasm, with a round nucleus and hyaline vesicle, but no distinct cell-wall (fig. 77, *c*). According to *Kovacs*,¹⁰⁶ the *Amœba coli* is a cause of enteritis, and has to do with the development of hepatic abscesses.

Similar organisms have been found in the intestine by *Lambl*.¹⁰⁷ *A. Schuberg*¹⁰⁸ has noticed that after purgatives have been administered, especially Carlsbad salts, amœbæ become very abundant in normal human faeces. *Kartulis*, *Massuitin*, *Osler*, *Dock*, *Quincke*, and *Roos*¹⁰⁹ have seen amœba-like organisms in the stools of patients suffering from dysentery and chronic enteritis.

2. **Sporozoa**.—Again reverting to *Leuckart's* classification, the sporozoa which chiefly concern us here are the oval psorospermia, the group to which belong the coccidia which infest the intestinal tract of man (*Dressler*, *Gubler*, *Kjellberg*, *Eimer*)¹¹⁰ and the liver (*Podivyssovskii*).¹¹¹ They appear in the faeces as elliptical forms, 0.022 mm. long, furnished with a thin membrane, and enclosing within their surface a number of granular nuclei, arranged for the most part in groups. They are to be found in great numbers. Their favourite seat is the intestinal mucous membrane, where they burrow, doing much damage to its structure. For this reason *Leuckart* appropriately names the parasite *Coccidium perforans*.

3. **Infusoria**.—1. *Cercomonas intestinalis*.—This protozoon was first found by *Lambl*¹¹² in the jelly-like mucous discharges of children, and it was afterwards observed by *Daraine*, *Marchand*, and *Zunker*.¹¹³ It is of pyriform shape, clearly nucleated, and furnished with eight tentacles of varying length (see fig. 78, *a*). *Davaine* discovered it in cholera, *Marchand* in typhoid, and *Zunker* in nine cases of diarrhoea. It would appear from these facts that the *Cercomonas intestinalis* is apt to thrive in a gut which is already diseased, and, when present, tends to cause

severe diarrhoea. The observations of Zunker especially strengthen this conclusion. According to *Grassi* and *Schewiakoff*,¹¹⁴ the parasite causes anaemia and diarrhoea in human beings, and by the changes which it effects in the mucous membrane interferes with absorption from the intestine. The Cercomonas was observed by *Müller*¹¹⁵ in the jejunum of a healthy man.

The *Megastoma entericum*, lately described by *Grassi*, is almost certainly identical with Lambi's Cercomonas.¹¹⁶ *Perroncito*¹¹⁷ has observed the encysted forms of the parasite in the intestine (fig. 78)—observations confirmed by the author in the case of children.¹¹⁸

Other Cercomonads have been seen in the same situation (*Davaine*), (see fig. 77).

2. *Trichomonas intestinalis*.—This is a pear-shaped organism, somewhat larger than Cercomonas intestinalis, and distinguished from it by bearing a ciliated disc at one extremity (fig. 77). It has been obtained from the intestine by *Marchand* and *Zunker*.¹¹⁹

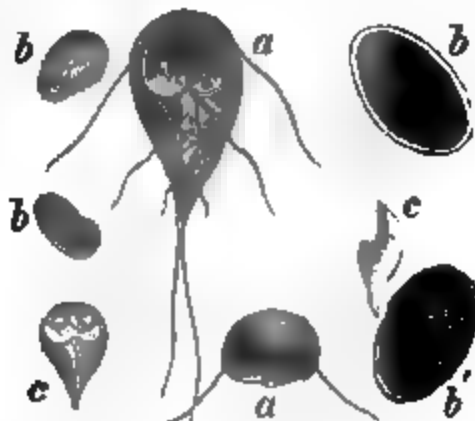


FIG. 78.—Cercomonads from the Stools.

- a. *Megastoma entericum* (*Grassi*).
- b, b'. Encysted forms of *Cercomonas intestinalis*.
- c. *Cercomonas intestinalis* after loss of its tentacles (*Lambi*).

3. *Paramaecium coli* (*Balantidium coli*).—An entozoon first found by *Malmsten*¹²⁰ in the discharges of diarrhoea, and afterwards observed by *Stieda*, *Graziadei*, *Perroncito*, and *K. Ortmann*.¹²¹ It is oval, 0.1 mm. long, and covered with cilia, which are planted more thickly around the buccal (?) orifice. The anterior extremity is smaller than the posterior, and the opening (anal) in this situation is but sparingly provided with cilia. The abdominal surface is less arched than the dorsal. Internally it is furnished with a nucleus and two contractile vesicles, and frequently contains amyloid particles and fatty granules. Its presence appears to be associated with diarrhoea. It also develops ulcerous processes in the colon, which induce peritonitis, and may thereby cause the death of the patient harbouring this entozoon (*Janowsky*,¹²² *Woit*¹²³).

In addition to the varieties mentioned here, other infusoria are occasionally present in the intestine (*v. Jaksch*).¹²⁴

2. Vermes.—The investigation of the faeces for intestinal worms has become of late years a matter of special interest to physicians, because experience daily teaches us that, even in temperate climates, the alimentary canal is apt to be beset with certain parasites of this class, which must be reckoned amongst the most formidable pests of mankind; and it often happens that an accurate diagnosis of their nature can alone enable the physician to adopt intelligent methods for their removal, and so, it may be, to save his patient's life.

Class I.—Platoda.

(a.) **Cestoda.**—The following tapeworms concern us here:

1. *Tænia solium*
2. *Tænia saginata* (*medicaneallata*).
3. *Tænia nana*.
4. *Tænia diminuta* (*flavopunctata*).
5. *Tænia curumerina* (*elliptica*).
6. *Bothriocephalus latus*.



FIG. 79.—*Tænia solium*: Head (magnified), Proglottis (actual size), and Egg (magnified). (Zeiss 3 eyo-piece IV, objective IV.) From a preparation by Cori and e. Jakob.

1. *Tænia solium*.—The *Tænia solium* may measure upwards of two or three metres. Its head is quadrilateral, about as large as a pin's-head ($\frac{1}{4}$ to $\frac{1}{5}$ of an inch), and dark in colour. This is succeeded by a delicate, thread-like neck, about an inch in length, and unjointed. The segments, or proglottides, which form the rest of the body, are short and relatively broad near the neck; but as they increase in size this relation ceases, and still growing in both dimensions, their quadrilateral form becomes evident about three feet from the head. Their average length is from 9 to 10 mm., and their breadth 6 or 7 mm.

Under the microscope the head is seen to present four prominent suckorial discs, usually pigmented, and between them a rounded elevation or rostellum, which is surrounded with about twenty-six hooklets of different sizes.

The sexual apparatus first becomes visible about a foot from the head (*Bristowæ*). The uterus is but little branched, and the genital pores are situated somewhat behind the middle of each segment.

The eggs are oval in shape, about 0.03 mm. in diameter and 0.036 mm. long, and surrounded with a thick shell, which is radially striated. When the eggs are mature, they may be seen to contain embryos furnished with hooklets (fig. 79).

In view of the possibility of infection with *Cysticercus cellulose* from the eggs of this worm, the greatest care is needed during the period of its destruction medicinally, both by the patient, who should be prevented from vomiting, and also by the medical and other attendants.

2. *Tænia saginata* (*mediocanellata*).—This parasite is longer than the *Tænia solium*, attaining to four or five metres, and its segments are also longer. The head is surrounded with four large and usually

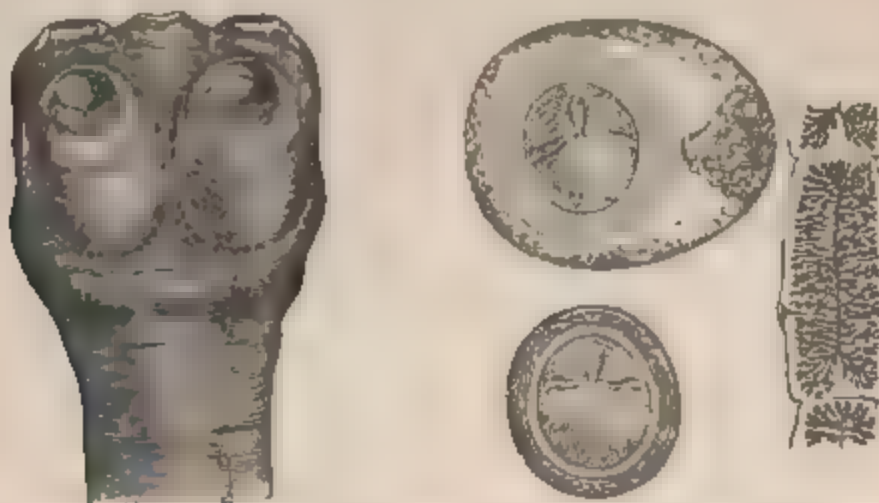


FIG. 80.—*Tænia saginata* (Hedw.). Proglottis, Egg. (Zeiss's eye piece II., objective IV.) From a preparation by Dr. Cor.

black pigmented suckers, but is not provided with a rostellum, and is without a circle of hooklets. The segments increase in length more gradually than in *Tænia solium*, and are commonly pigmented.

The uterus is very much branched, and the genital pore is situated at the side of the proglottis. The eggs resemble those of *Tænia solium*, but are more oblong, and exhibit the primordial yolk membrane (fig. 80).

3. *Tænia nana*. This parasite averages from 2.5 to 10 mm. in length, and its greatest breadth is about 0.7 mm. It has a globular head 0.3 mm. in diameter, furnished with four circular suckers, and a rostellum 0.06 mm. long, carrying twenty two to thirty hooklets at its anterior extremity, which is rounded off. The rostellum can be protruded to a considerable distance from the head or entirely withdrawn within it. In the latter position it has the form of an hour-glass. The body is attenuated in its anterior third, but proceeding backwards grows quickly

in bulk. The segments are short, and towards the end of the body are scarcely one-fourth so long as they are broad. The uterus is oblong and loaded with eggs of 0.03 to 0.04 mm in diameter. The shell does not exhibit the rod-like structure, but consists of a double membrane, within which are a spiral thread and amorphous matter, in which granules are embedded. In the interior of the egg may be seen the embryo, already provided with five or six hooklets (fig. 81). The parasite may be present in great numbers in the intestine, and is apt to produce severe nervous symptoms, such as epileptic seizures, insensibility and mental derangement, and melancholia (*Grassi, Comini*).¹²⁵

Recent observations go to prove that the *Tænia nana* is very widely distributed. It especially attacks children and young persons. It was first discovered by *Bilharz* in Egypt, and since then has been recognised as of common occurrence in Italy and Sicily.¹²⁶ It is probable that it

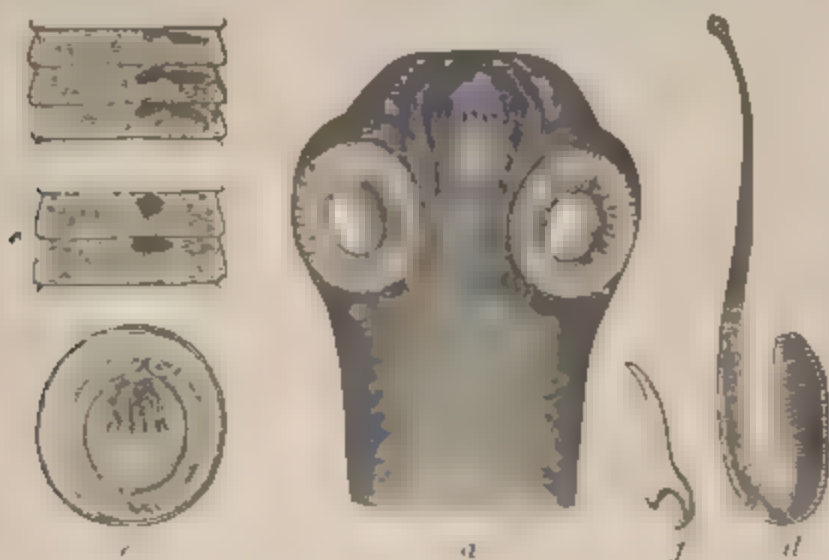


FIG. 81. *Tænia nana*. a. Head (with Rostellum drawn in). b. Immature Proglottis. c. Mature Proglottis. d. Animal (natural size). e. Egg. f. Hooklets. (Zeiss's eye-piece IV., objective VIII.) From a preparation by Dr. Comi.

exists elsewhere more frequently than is supposed. *Ranson* has published an account which goes to prove that it was observed in England so long ago as 1856.¹²⁷ According to *Grassi*, *Tænia nana* is in Sicily the commonest parasite. The number present in a single individual may reach 4000–5000. *Mertens* has shown that it occurs in Germany.¹²⁸

4. *Tænia diminuta seu flavopunctata*. — *Weinand*, and after him *Leroy* and *Perrone*,¹²⁹ have described a parasite of the human intestine which they called *Tænia flavopunctata*. It was later identified by *Grassi* with *Tænia diminuta*, which infests certain of the rats. It is allied to *Tænia nana*, is 20–60 cm. in length, has a rudimentary rostellum and no hooklets. The eggs are twice as large as those of *Tænia nana*. The embryonal integument is thickened at the poles.

5. *Tænia cucumerina (elliptica)*. This is a worm which averages 15–50 cm. in length. The head is furnished with four suckers and a

prominent rostellum upon which the hooklets are arranged in five or six rows. The first forty proglottides are quadrilateral, those which succeed elongated and rounded, and measure at the extremity 6 15 mm. by 3 mm. in breadth. The mature segments are of a reddish colour, and readily separate from those nearer to the head. The parasite is distin-

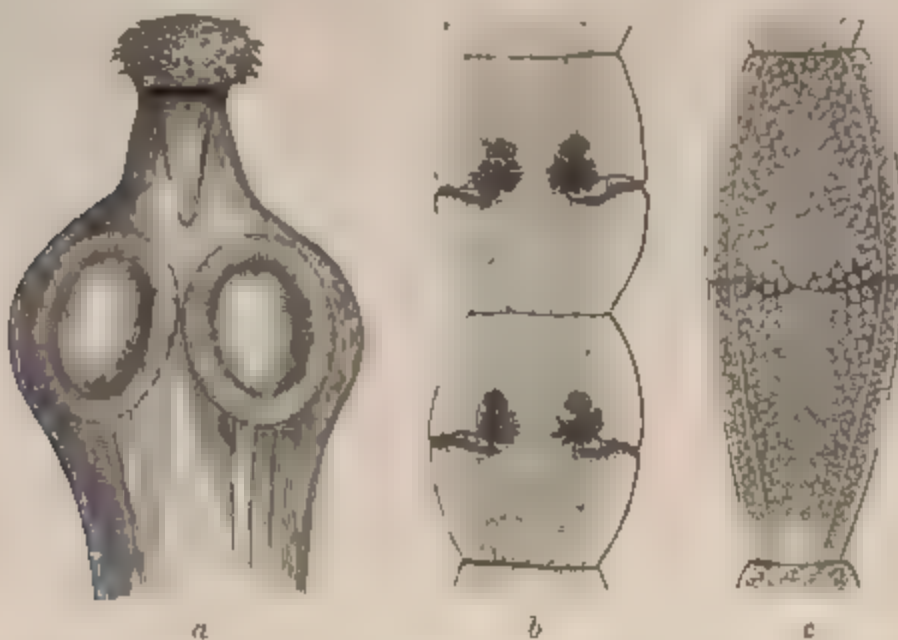


FIG. 82.—*Taenia cucumerina*. *a*, Head. Proglottis (magnified). *b*, Immature. *c*, Mature (Zeiss's eye-piece II., objective a*.) After Dr. Cori.

guished from others which infest man by the twofold uterus and genital pore in each segment. The eggs, when shed, have a diameter of 0.05 mm., and contain an embryo furnished with hooklets.

The parasite is not at all rare amongst human beings;¹³⁰ it especially

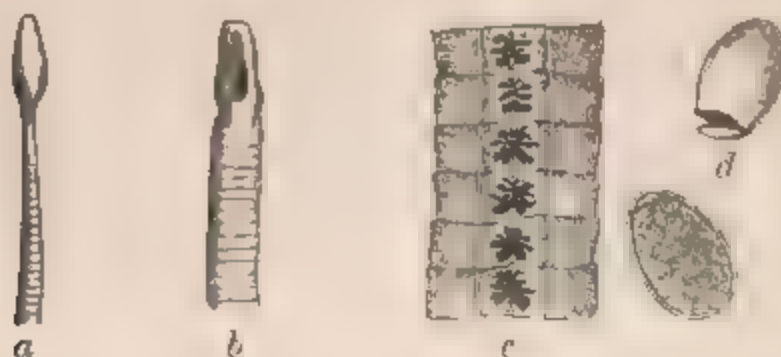


FIG. 83.—Head of *Bothriocephalus latius* (*a*, eye-piece III., objective IV., Reichert; *d*, Zeiss's eye-piece VI., objective IV.). After Dr. Cori.
a, Seen on edge; *b*, Seen on the flat; *c*, Proglottides; *d*, Eggs.

infests children, and is acquired by them from dogs by swallowing the dog-louse (*Trichodectes canis*), which is the host of the larva.

It is sufficient merely to mention here the rarer forms of tapeworm known to be present in the human intestine, such as *Taenia madagascariensis* (Grenel).¹³¹

6. *Bothriocephalus latus*.¹³²—This worm attains a length of 5–8 metres. The head is ovoid, and 2 mm. long by 1 mm. broad. It is cleft, and provided with two lateral suckers, placed on either side of the middle line. It has no hooklets. The proglottides are at first short and small. They increase in breadth as they proceed, and towards the end of the parasite approach the square form.

The uterus of mature proglottides containing eggs exhibits a retiform arrangement, and appears superficially as a small rosette. This uterine rosette is characteristic of *Bothriocephalus latus*.

The eggs are oval, and measure 0.07 mm. by 0.045 mm. They are covered with a brown shell, and open by a small lid at one end. They contain centrally transparent masses of protoplasm of uniform size. The arrangement of the genital organs makes it certain that they will be shed, and thus affords a secure resource to diagnosis. *Bothriocephalus latus* has recently acquired an increased clinical significance, inasmuch as it has been found frequently by different observers in conjunction with the symptoms of pernicious anaemia.¹³³

Other varieties of this parasite have been observed, as the *B. cordatus* in Greenland,¹³⁴ and the *B. liguloides* in Japan and China. The latter infests the sub-peritoneal tissue, and especially that of the lumbar region in man.¹³⁵

The presence of one of the tapeworms here described is made evident —apart from clinical symptoms, which do not concern us here—by the discovery of proglottides in the stools.

A careful microscopical examination with a medium power (Hartnack objective IV., Zeiss objective C., Reichert objective IV.) may show the eggs in the faeces. When it is supposed that a patient is suffering from tapeworm, whilst yet no evidence of the parasite can be derived from a careful inspection of the stools, it is well to mix these with water, which is constantly poured off and renewed until the greater part of the faecal mass has been dissolved. An examination of the sediment will then probably reveal the eggs, if the suspicion be well founded. Stenbeck's sedimentator may be used with advantage here. It is only in the case of *Bothriocephalus*, however, that eggs are sure to exist in the faeces. This is not so with the tæniae. To determine to what form of tapeworm a particular proglottis belongs, the best proceeding is to examine the specimen, mounted in glycerine, with a low magnifying power; and the same thing may be done if a head be obtained.¹³⁶

Cysts of echinococcus and hooklets may be found in the faeces when a hydatid cyst has burst into the intestine. Heller¹³⁷ was able by the discovery of such a cyst to diagnose a doubtful liver complaint as one of hydatids.

(b.) **Trematoda.**¹³⁸—The varieties of distomata, *D. hepaticum* and *D. lanceolatum*, have been found in rare instances in the intestine or biliary passages of man.

1. *Distoma hepaticum*.—This is a leaf-shaped worm, measuring 30 mm. by 12 mm. The head is short, and furnished with a sucker. There is another sucker on the ventral aspect, and in front of this the genital pore is situated. The latter leads to a uterus, which is convoluted like a ball of wool and crowded with eggs. On the posterior surface of the fore part of the body are a number of prickly scales. By their means the parasite can be recognised from a fragment derived from the liver or fæces. The alimentary canal is bipartite, and often distended with blood and bile.

The eggs are oval, 0.13 mm. long and 0.08 mm. broad, brown in

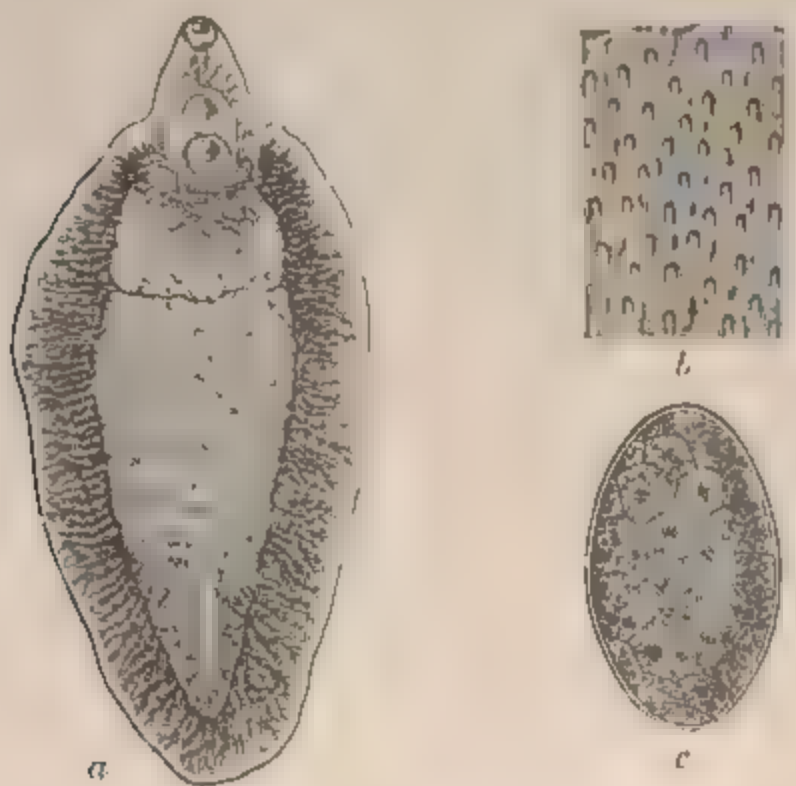


FIG. 84.—*Distoma hepaticum*.

a Animal; b. Skin with prickly scales, c. Egg (b. Zeiss x eye-piece II, objective ap. 0.1
c. Zeiss x eye-piece II objective IV.). From preparation by Dr. Cor-

colour, and covered with a shell consisting of two layers; one end is broader than the other, and opens by a small lid, like the egg of *Bothriocephalus* (see fig. 84), which, however, is not so brown in colour. They contain daughter cells, which are visible through the shell. The worm enters the intestine and is voided with the fæces. *Biermer*, *Bostroem*, and *Baelz*¹³⁹ have described these parasites as occurring in human beings.

2. *Distoma lanceolatum*.—This entozoon is from 8 to 10 mm. long and 2 to 3.5 mm. broad. It is lance-shaped, as its name implies, and pointed at either extremity, but more so in front than behind. The surface is without scales, and the intestine bipartite but unbranched (see

fig. 85). Behind the ventral sucker are two lobular testicles, and behind these the convoluted uterus filled with small brown eggs.

The eggs are 0.04 mm. long, and 0.03 mm. broad, and they contain the

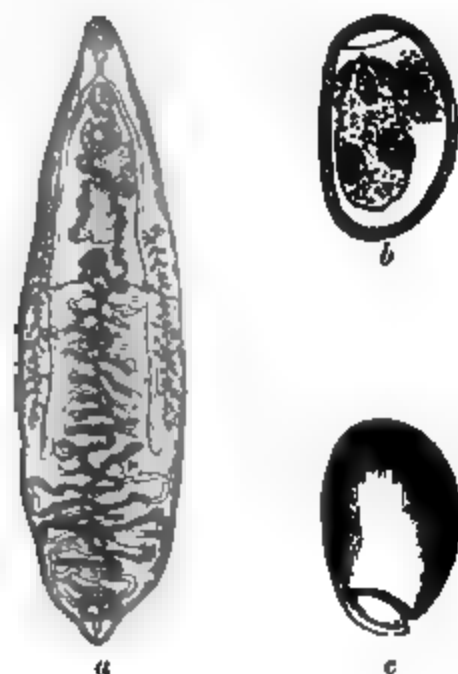


FIG. 85.—*Distoma lanceolatum* (seven times actual size) and Eggs: a. With embryo; b. Empty shell (Zeiss's compensation eye-piece IV., objective IV.). From preparation by Dr. Corr.

mature embryo. *Bizzozero*¹⁴⁰ maintains that when the worm is present in the intestine its eggs are to be found in the stools. The observations of *Baelz*¹⁴¹ corroborate this view. *Perroncito*¹⁴² found the eggs of this fluke in the case of persons infested by *ankylostoma*.

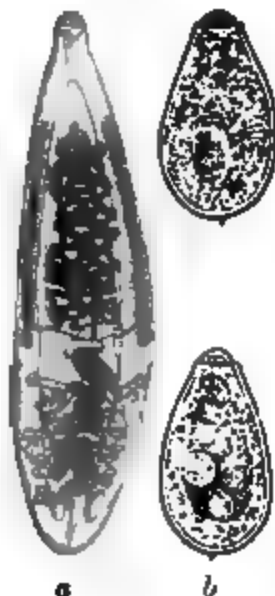


FIG. 86.—*Distoma sinense* (*spathulatum*). a. Animal; b. Eggs (Zeiss's compensation eye-piece, objective IV.). From preparation by Dr. Corr.

The last two forms of parasites but rarely occur in man, and consequently neither they nor their ova are often to be seen in the faeces. Moreover, they very seldom cause serious symptoms.

3. *Distoma rathonisi*.¹⁴³—This distoma was described by *Rathonis*, who observed the first specimen in a Chinese woman, thirty-seven years of age. The host suffered from severe pains, referred to the liver. The parasite resembles *D. hepaticum*, but is larger (25 mm.); the ramifications of the alimentary canal are less complex, especially towards the hinder end.

For further particulars *Piorier* may be consulted.

4. *Distoma sinense* (*D. spathulatum*).—This parasite, about 18 mm. long, is, when living, of a reddish colour, and its covering is so transparent that the viscera are clearly visible. It is broad in the middle, and pointed at either end, but more so anteriorly (fig. 86). The buccal is considerably larger than the abdominal sucker. Behind the latter is a simply convoluted uterus, and behind this two testicles having a star-

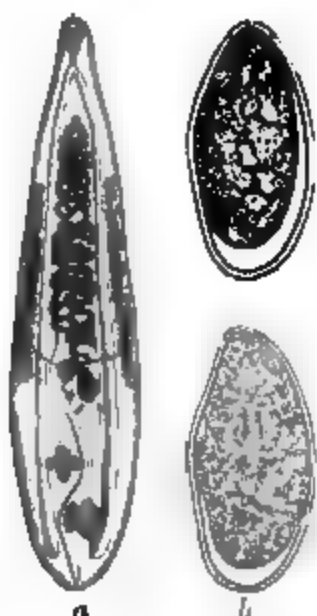


FIG. 87.—*Distoma felineum* (*sibiricum*): a. Animal; b. Eggs (Zeiss's compensation eye-piece, objective IV.). From preparation by Dr. Cori.

shaped lobular arrangement. The alimentary canal is bipartite and unbranched. The eggs are oval, lidded, and spiked at the opposite end (fig. 86). They are 0.028 to 0.3 mm. long, and 0.016 to 0.017 mm. broad. The parasite is endemic in parts of Japan, and gives rise to serious liver trouble.

5. *Distoma felineum* (*D. sibiricum*, *D. Winogradoffi*).¹⁴⁴—This fluke resembles the last in shape and appearance. The chief difference is in the testicles, which in *Distoma felineum* consists of 4 to 5 smooth lobes, which are not further divided (fig. 87). The eggs are 0.026 to 0.03 mm. long, and 0.011 to 0.015 mm. broad, lidded at the pointed end; more oblong than those of *D. sinense*, and somewhat flattened on one side. The usual host of this parasite is the cat or the dog, but it has lately been observed in the human inhabitants of Tomsk by *Winogradoff*.

CLASS II.—ANNELIDA.

i. Order Nematoda (*Round Worms*).

a. Family Ascaridæ.¹⁴⁵

1. *Ascaris lumbricoides* (common round worm).—This is a cylindrical worm, of some size, with a body that tapers from before backwards. The male is 250 mm. and the female 400 mm. long. The head, which is distinct from the body, consists of three conical prominences (lips) furnished with tactile papillæ and minute teeth. The caudal process of the male is folded hook-like on the abdominal surface, and is provided



FIG. 88.—*Ascaris lumbricoides*.

a. Head; b. Hind sexual end of male; c. Egg; d. Male.

b. Slightly magnified; c. (eye-piece I., objective 8A, Reckert); d. half natural size.

with papillæ. In the female the vulva lies deeply behind the anterior third of the body (fig. 88). The eggs are nearly round, and brownish yellow in colour. Their diameter is 0.06 to 0.07 mm. In the fresh state they are covered externally with an albuminous layer, and beneath this is a tough shell, which in turn encloses the very granular contents.

The *Ascaris lumbricoides* infests the small intestine in man, and it appears to be common to all climates. It occurs also in cattle and in sheep. It has no special medical interest; but it is thought by Lutz¹⁴⁶

to cause spasm and tympanitis, and to impede nutrition in children. *Kartulis*¹⁴⁷ records a case of death in a man following directly upon an invasion of the liver by ascarides. Severe nervous symptoms also, as amaurosis, strabismus, and evidence of meningitis,¹⁴⁸ have been found to accompany their presence in great abundance. They have caused death by invading the liver.¹⁴⁹

2. **Ascaris mystax** (round worm of the cat).—This worm closely resembles the preceding, but is smaller, and is readily distinguished from it by the shape of its head, which is pointed, and bears two lateral wing-shaped processes. The male is 45–60 mm. long and the female 110–120 mm. The eggs are globular, larger than those of *A. lumbricoides*, and covered with a dimpled shell (fig. 89).

3. **Oxyuris vermicularis** (common thread-worm or teat-worm).—The

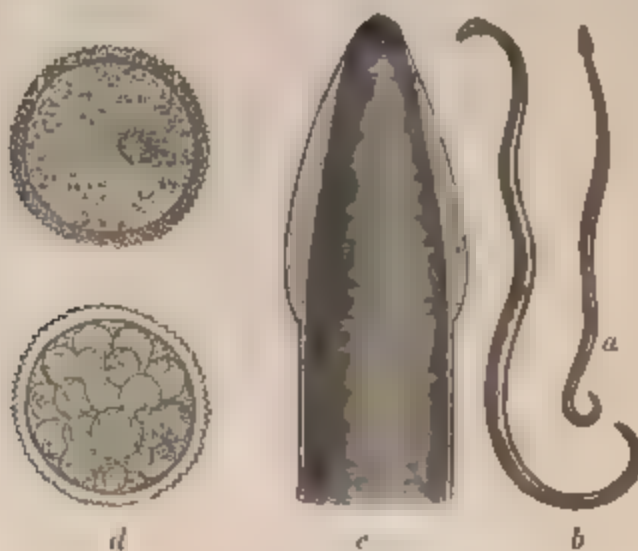


FIG. 89.—*Ascaris mystax.*
a. Male; b. Female; c. Head; d. Egg.
a, b. Half natural size; c. (eye-piece II., objective IV., Zeiss); d. (eye-piece IV., objective VII., Zeiss). After Dr. Cort.

female is 10 mm. in length, and exhibits two fully-developed uteri, which extend symmetrically backwards from the vaginal orifice. The male is rather less than half the length of the female, and its tail is provided with six pairs of papillæ. The head of both sexes is similar. It displays a remarkable cuticular enlargement and small prominent lips.

The eggs are irregularly oval, and measure $\frac{1}{15}$ (0.05 mm.) by $\frac{1}{100}$ (0.02–0.03 mm.) inch. The shell is membranous, and consists of two or three laminæ. Its contents are coarsely granular. The eggs often contain an embryo with an indistinct alimentary canal and a tail equal to half the entire length. The presence of the parasite is attended with uncomfortable sensations, among which itching in the situation of the anus is prominent.¹⁵⁰

β. *Family Strongylidae (Leuckart).*¹⁵¹—To this family belongs one

of the most important and formidable of the parasites which infest the human intestine. This is—

Anchyllostoma duodenale (*Dochmias duodenalis*, *Strongylus duodenalis*).—It was formerly believed that this worm occurred only in the tropics and in certain districts of Italy.¹⁵² Of late years, numerous observations in Egypt (*Sandwith*), Italy (*Perroncito* and others¹⁵³), Germany, Switzerland (*Menche* and others¹⁵⁴), and Belgium¹⁵⁵ have made it clear, together with the older researches,¹⁵⁶ that the inhabitants of temperate regions are not free from its attacks.¹⁵⁷

The Anchyllostoma is cylindrical in form. The male measures 8 to 12 mm. in length, the female 10 to 18 mm. The anterior extremity

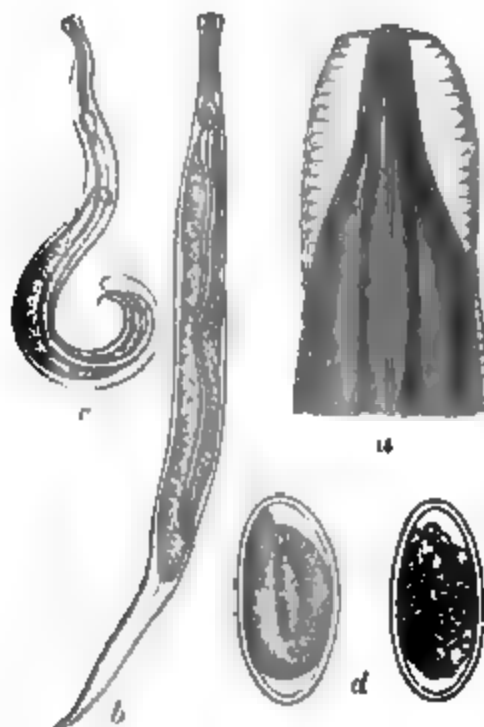


FIG. 90.—*Oxyuris vermicularis*.

a. Head ; b. Female ; c. Male ; d. Eggs.

a. (eye-piece II., objective IV., Zeiss); b. c. slightly magnified; d. (eye-piece IV., objective VII., Zeiss). After Dr. Cori.

is pointed, and reflected towards the dorsal surface. The oral orifice is armed with four claw-like teeth. The caudal extremity of the male expands into a pouch with three flaps; that of the female is pointed and conical; the vulva is situated behind the middle third of the body.

The eggs are smooth and oval, measure 0.05 to 0.06 mm. in length and 0.03 to 0.04 mm. in breadth, and usually contain two or three large daughter-cells. The embryos develop rapidly outside the human body. In stools which contain eggs, the embryo may be seen and observed after the lapse of twenty-four to forty-eight hours.

Except when anthelmintics have been administered, the eggs are the only signs of the parasite to be found in the discharges, and it is con-

sequently of the utmost importance to be familiar with their appearance. The accompanying figure shows this at various stages of their development (fig. 91).*

The presence of *Anchyllostoma* is to be suspected in all cases where a severe form of anaemia occurs in labourers (and especially in brick-burners, miners, and tunnel-borers), without any obvious and sufficient cause. It is, however, necessary to bear in mind that *Bothriocephalus latus* produces similar symptoms (p. 221). Examination of the stools will then afford the means of diagnosis. If the intestines contain *Anchyllostomia*, the characteristic eggs with their large blastomeres will

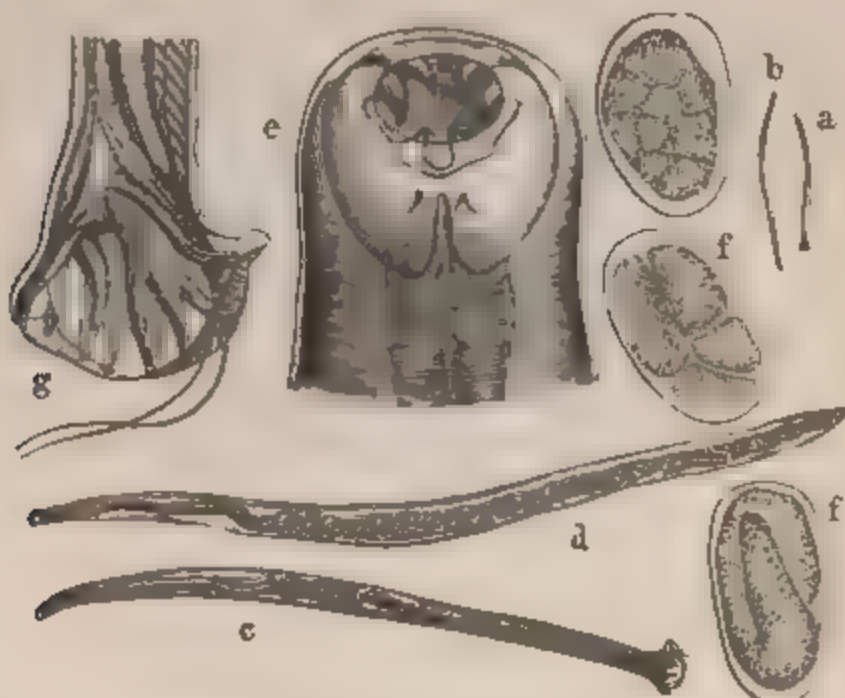


FIG. 91. *Anchyllostoma daedaleum*

- a. Male (natural size).
- b. Female (natural size).
- c. Male (magnified).
- d. Female (magnified).

- e. Head (eye-piece II, objective ap. 6.3 Zeiss).
- f. Eggs (eye-piece II, objective ap. 6.3 Zeiss) after Dr. C. J.
- g. Posterior end of male (eye-piece IV, objective V, Zeiss) after Prof. Chiaro.

be seen. If any uncertainty as to their nature should persist, the faecal substances containing them should be allowed to remain for some days in a warm place, and again examined microscopically. The process of segmentation of the egg will then be seen to have advanced, and fully developed embryos will be visible here and there. The administration of anthelmintics, and especially of the æthereal extract of male-fern, will cause the expulsion of the mature worm, and an inspection of the resulting discharges cannot fail to establish the diagnosis. [Sandwith¹ finds thymol by far the most efficient vermicide in cases of anchyllostomiasis.]

The character of the stools in this affection varies greatly. Diarrhoea

* [An instructive case of this disease is reported in the *Lancet*, February 1, 1890.]

is usually present, and blood is frequently passed. But it may happen that the discharges are altogether normal. They have been known to contain great quantities of Charcot-Leyden crystals (*Leichtenstern*, see *ante*). Investigations by *Bohland*¹⁵⁹ on the subject of metabolism in men infested by *Anchylostoma*, point to the conclusion that the increased distribution of proteids is due to a toxine secreted by the parasite. [In the report of the Ceylon Commission, 1887,¹⁶⁰ *ancylostomiasis* was stated to be a cause of beri-beri. The association is probably accidental (*Sonsino*).]¹⁶¹

γ. Family *Trichotrachelidae* (*Leuckart*).¹⁶²

1. **Trichocephalus dispar** (whip-worm).—This worm has a whip-like form, consisting of a short, stout hinder part, and a long, spiral, filiform

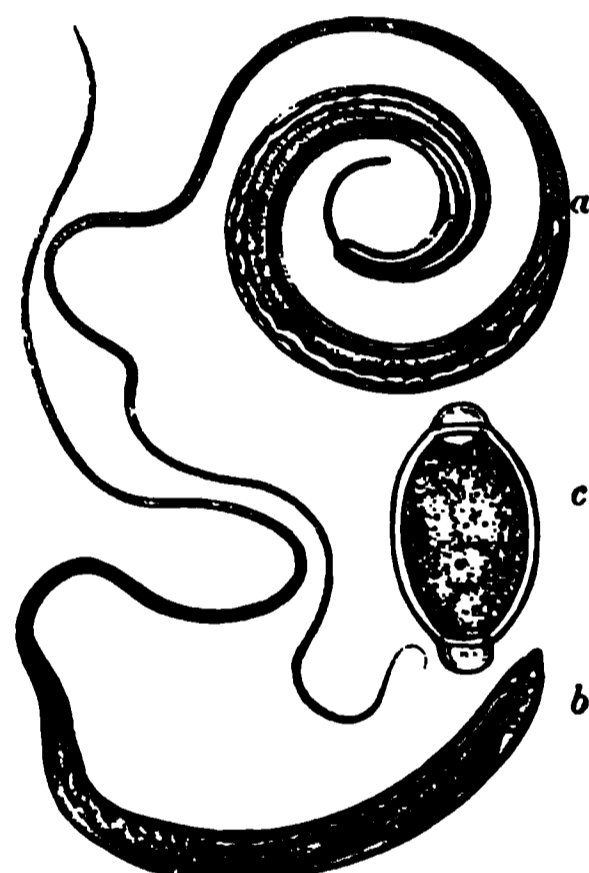


FIG. 92.—*Trichocephalus dispar*.

a. Male; b. Female; c. Eggs; a. b. slightly magnified; c. (eye-piece II., objective IV.. Zeiss). After Dr. Cori.

process anteriorly. The male is 40 mm. and the female 50 mm. long. The short hinder part measures 1 mm. in thickness (fig. 92).

The eggs, which are occasionally to be found in the faeces, are brown in colour, 0.05–0.06 mm. long and 0.02 mm. broad. The shell shows a double contour, and is flattened at either end, where it is furnished with a small lid, formed of a glossy substance. The yolk is very granular (fig. 92). According to *Erni*¹⁶³ and others, this parasite, together with *Anchylostoma* and an insect-larva, produces the beri-beri disease which is endemic in Sumatra. This view is, however, opposed by other writers (*Scheube*, *Scheffer*).

2. **Trichina spiralis**.¹⁶⁴—Trichina occurs in two different forms in

the human body, according as its habitat is the muscular tissue or the intestine. It is with the trichina of the intestine that we have to do here, since this form alone is found, though rarely, in the faeces. The male is 1.5 mm. in length, and the female 3 mm. The former has four prominent papillæ situated between the conical protuberances at the extremity. The sexual organs of the female consist of a tubular ovary, which is placed at the hinder part of the body, and a tubular uterus, with which the ovary communicates in front (fig. 93). Impregnation takes place in the intestine. The eggs develop into embryos while still

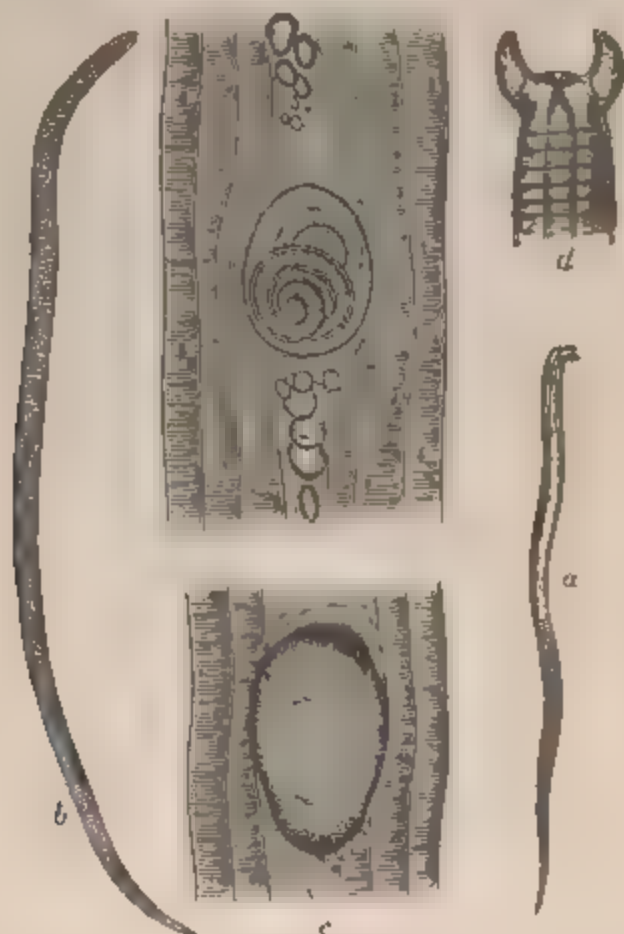


FIG. 93. Trichinæ.

a. Male, and b. Female Intestinal Trichinæ (slightly magnified), c. Trichinæ of muscle; d. Posterior end of male (σ . h. eye-piece II, objective IV; c. eye-piece IV, objective IV, Zeiss).

within the uterus, and the newly born parasite almost immediately perforates the gut, and becomes embedded in the muscles of its host.

'Of their own accord these worms rarely pass with the stools. But in any case where there is reason to believe that trichinous meat has been eaten and trichinosis is apprehended, an anthelmintic may be given, and the detection of intestinal trichina in the faeces will establish the diagnosis in a very early stage of the affection. As recently shown, however, in a case published in the author's clinic by v. Strausly,¹⁶⁵ an accurate diagnosis can only be made by excising a portion of muscle.'

Researches made by *Th. K. Brown*¹⁶⁶ show that an increase takes place in the number of eosinophil cells present in the blood. *Brown* believes that this indication may be utilised in diagnosis.

d. *Rhabdonema strongyloides* (*Leuckart*).—Certain nematode worms have been discovered in the stools in cases of Cochin-China diarrhoea (*Normann, Bayay, Seifert*).¹⁶⁷ They occur commonly in conjunction with *Anchylostoma* (*Grassi, Parrona, Perroncito*).¹⁶⁸ It was formerly thought that there were two distinct parasites of this kind, viz., *Anguillula intestinalis* and *Anguillula stercoralis*; but more recent observations by *Leuckart* and *Grassi*¹⁶⁹ have shown that the latter is only an intermediate form in the development of *A. intestinalis*. A complete

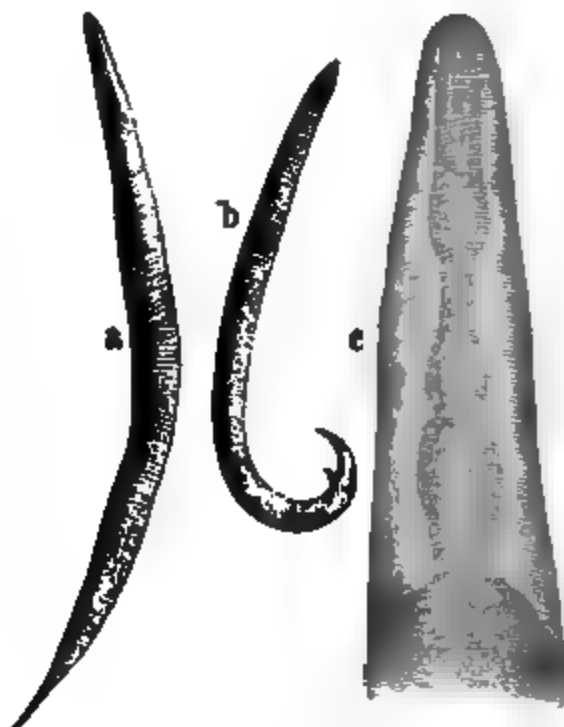


FIG. 94.—*Anguillula stercoralis*.

a. Female; b. Male; c. Head (eye-piece II., objective 8A, *Reichert*).

knowledge of the subject, however, implies an acquaintance with this form, as it may be confounded with other *Helminthidae*. The life-history, according to *Grassi*, is as follows:—

The *Anguillula* which infests the human intestine deposits eggs, from which the young are produced as embryos or larvae. These are discharged with the stools. As found there, they already exhibit sexual maturity (*Anguillula stercoralis*) and produce embryos, which undergo no further change in the human system. The body of this worm is round, and shows faint traces of transverse striation. The head is in the form of a blunted cone (fig. 94) and sessile upon the body. It is furnished with two lateral jaws, each bearing a pair of teeth. The male is 0.88 and the female 1.22 mm. long. *Anguillula intestinalis*

measures 2.25 mm. in length and 0.04 mm. in thickness at its middle. It has a triangular mouth closed by three small lips. The vulva lies at the junction of the middle with the posterior third. Its habitat is the small intestine. The eggs resemble those of *Anchylostoma duodenale*, but are longer, more elliptical, and pointed at the poles. In recent stools the larvæ alone are to be seen. It is doubtful whether the parasite is directly mischievous; but its constant association with *Anchylostoma*, and the readiness with which the two may be confounded, render its recognition a matter of consequence.

3. Insects.—It requires to be noticed that insect larvæ infest the stools. *Joseph*¹⁷⁰ has reported a number of species, which are for the most part taken into the intestine with food (cheese, meat), and give rise to a variety of morbid symptoms, as colicky pains, vomiting, &c. Special mention may be made of the immature cheese-maggot (*Piophila casei*) and *Drosophila melanogaster*, which are derived

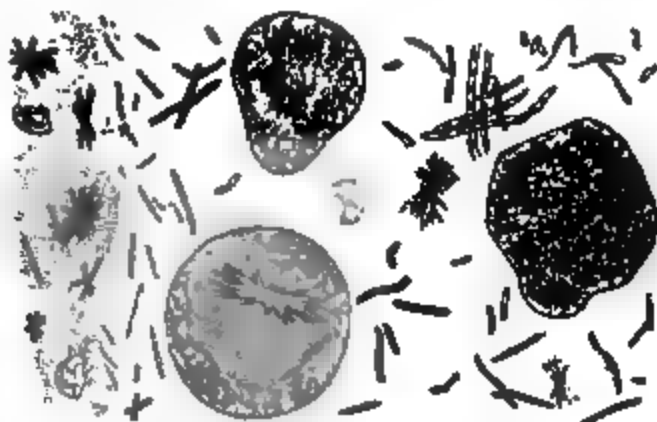


FIG. 95.—Hematoidin Crystals from Acholic Stools (eye-piece III., objective 8A, Reichert).

from curdled milk, and may attain the chrysalis form in the intestine before they are discharged per rectum. Of other species are the larvæ of three varieties of *Homalomyia*, of *Hydrothæa meteorica*, *Cyrtoneura stabulans*, *Calliphora erythrocephala*, *Pollenia rudis*, *Lucilia cæsar* and *regina*, *Sarcophaga haemorrhoidalis* and *haematodes*, and *Eristalis arbustorum*, all of which are apt to occur. *Rembold*, *Lampa*, and *Kohn*¹⁷¹ have also observed in the stools certain lancet-shaped bodies, 8 mm. long, which are covered with hair and indented on the surface, and these have been identified by *v. Graff* as the larvæ of *Anthomyiæ*. [*Finlayson*¹⁷² records a case in which swarms of larvæ were passed alive from the bowel of a man. He identified the insect as *Anthomyia canicularis* or *scalaris*.] *Henschen*¹⁷³ records a whole series of analogous observations.

4. Crystals.—Crystalline bodies are a common constituent of the fæces, and in some cases are to be found in great quantities. They may be organic or inorganic.

1. Charcot-Leyden Crystals.—These bodies, of which an adequate description has already been given in connection with the *Blood* (Chap. I.) and the *Sputum* (Chap. II. and fig. 64), are comparatively rarely to be seen in the faeces. *Nothnagel* has met with them in typhoid fever, and *Leichtenstern*¹⁷⁴ in phthisis and ancylostomiasis. Their presence, however, has no pathological significance.

2. Hæmatoidin Crystals.—It is surprising that so little attention has been paid to this subject. *Uffelmann*,¹⁷⁵ indeed, remarks that crystals of hæmatoidin are sometimes present in the discharges of infants at the breast; but apart from this notice the matter has been passed by in silence. The author has found these crystals often enough in the faeces, especially in chronic intestinal catarrh from over-eating, and in many instances in which blood had been discharged into the intestine some time (several days) before the stools were passed. They usually exhibit an ill-marked crystalline structure. Particularly good specimens were

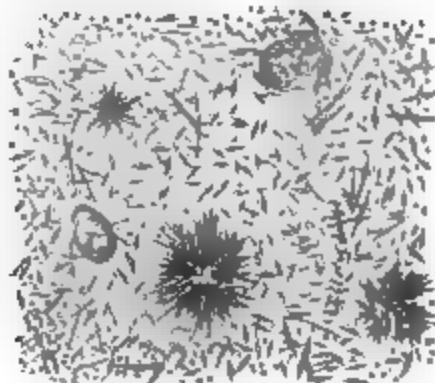


FIG. 96.—Acholic Stools (eye-piece III., objective $\frac{1}{3}$, oil immersion, *Reichert*; *Abbe's* mirror, narrow diaphragm).

seen in the stools of a patient suffering from nephritis. The crystals are sometimes free, and sometimes enclosed in masses of a shining substance which resembles mucin (fig. 95). Similar crystals were found in the liquid motions of a man who suffered from pernicious anaemia (*v. Jakob*).

3. Cholesterin.—This substance enters normally into the constitution of the faeces, and can always be obtained from them.

It seldom appears in its crystalline form (fig. 130), according to *Nothnagel*, and the statement is undoubtedly correct. (For the microscopical and chemical character of cholesterin, see chapters on *Sputum* and *Urine*.)

4. Fat Crystals.—*Nothnagel*, in his well-known monograph,¹⁷⁶ states that fatty substances occur in the faeces in the form of needles. *Gerhardt*¹⁷⁷ found quite an enormous quantity of organic crystals in the stools of jaundice (fig. 96). He was himself of opinion that such bodies

consisted of tyrosin; but one of his pupils, named *Oesterlein*,¹⁷⁸ who pursued the investigation further under *Gerhardt's* auspices, came to the conclusion that chemically they were lime and magnesia salts of the higher fatty acids, and that, therefore, soaps of lime and magnesia bases form a part of such discharges. According to *Stadelmann*,¹⁷⁹ they are sodium soaps. The author, however, can add his own testimony to *Gerhardt's* statement, for he has repeatedly seen great quantities of crystals arranged in clusters in acholic stools; and is disposed, on the grounds of separate investigation, and especially of the ascertained character of similar bodies occurring in other excretions (see chapter on *Urine*), to believe that such crystals do not consist of tyrosin, but rather of a combination of the alkaline earths with the higher fatty acids.

Uffelmann,¹⁸⁰ who had already observed these crystals in the discharges of infants, arrived independently at the conclusion that they could not consist of tyrosin. According to *Fr. Müller*,¹⁸¹ their presence indicates an impediment to the absorption of fat from the intestines.

In cases of jaundice, so frequent among children, such crystals are profusely present in the stools, and they are normally found in those of infants during lactation (*v. Jaksch*).

5. Oxalate and other Organic Salts of Lime.—Oxalate of lime is a sufficiently common constituent of the fæces, and appears on microscopical examination (see fig. 118). According to *Nothnagel*, it is always derived from the food. It is most abundant after a vegetable diet, and when manifested in considerable quantity the discharges also contain abundant débris of plant-tissues.

*Uffelmann*¹⁸² asserts that crystals of lactate of lime occur in the discharges of children as sheaves of radiating needles. Other organic salts of lime, as the acetic and butyric acid salts, have been observed in the discharges of persons suffering from acute gastric and intestinal catarrh (*v. Jaksch*).¹⁸³

6. Carbonate of Lime occurs rarely in the stools as amorphous particles and dumb-bell figures (fig. 131).

7. Sulphate of Calcium.—This salt is very seldom present in the stools. It can be obtained, however, by the action of sulphuric acid on the fæces, which shows that other lime salts are present. Its form presents the same variety here as in the urine (fig. 122).

8. Phosphate of Lime.—This substance crystallises in stout or elongated wedges, grouped together so as to form larger or smaller gland-like masses (fig. 121). The presence of such crystals in the fæces is without pathological significance.

Other salts of lime are occasionally met with. They are impregnated with bile-colouring matters, and deeply stained a yellow colour.

9. Triple Phosphate.—The phosphate of ammonia and magnesia

occurs sometimes as well-formed coffin-lid crystals (fig. 119), sometimes ill-defined crystalline masses, and very rarely in the elder-leaf arrangement (fig. 127).

Perfect crystals are most commonly to be found in fluid motions, and in the mucus which adheres sometimes to the faeces, whether liquid or solid. Fragments of coffin-lid crystals alone may be visible, and these often exhibit fissures and flaws, or there may only be mere splinters (*Nothnagel*). It is notable that these crystals seldom take up bile pigment. By their chemical constitution they may be readily recognised; and they dissolve easily, as has been already mentioned, in acetic acid (p. 137).

10. Sulphide of Bismuth Crystals.—When preparations of bismuth have been taken internally, the stools may be found to contain crystals

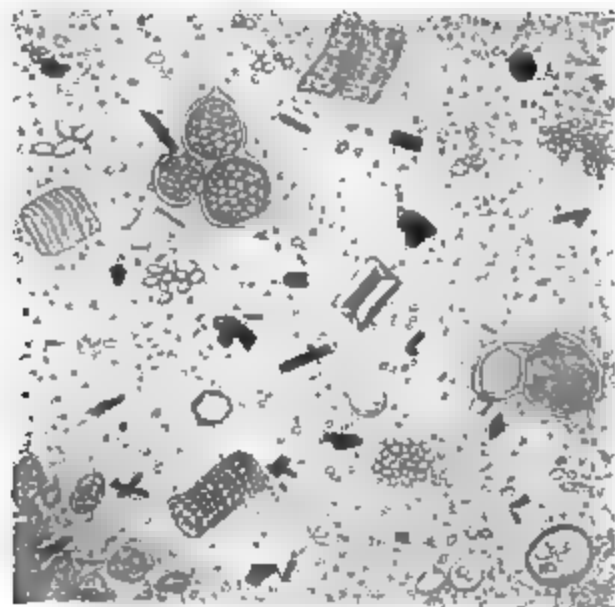


FIG. 97.—Sulphide of Bismuth Crystals from the Stools
(eye-piece III., objective 8A, *Reichert*).

which bear a remarkable resemblance to those of chloride of haematin (haemin). They consist of sulphide of bismuth, as may readily be shown by a comparison with the bodies formed when nitrate of bismuth is added to ammonium sulphide.

III. CHEMICAL EXAMINATION OF THE FAECES.—In striking contrast with the valuable information to be derived from the naked-eye and microscopic investigation of the faeces, is the very slight assistance which the chemical examination of those discharges lends to the purposes of diagnosis.

A. Organic Substances.

1. Mucin.—Mucin, as *Hoppe-Seyler*¹⁸⁴ has said, may be looked upon as the basis of the faecal substances, a statement which can be confirmed

by the author. *Fr. Müller*, on the other hand, asserts that mucin is not so abundantly present in the stools. For the detection of mucin in the stools the following process may be recommended :—The fæces are stirred up with water, their own bulk of lime-water added, and the mixture allowed to stand for some hours. It is then filtered, and the filtrate tested with acetic acid for mucin (see chapter on *Urine*).

2. Albumin.—The presence of albumin may be shown as follows :—The fæces are mixed with a large quantity of water, to which a trace of acetic acid has been added, and a watery extract is thus made. This is repeatedly filtered, and the filtrate may be submitted to the tests for albumin mentioned in the chapter on *Urine* (*q.v.*). Under ordinary circumstances no proteid reactions can be obtained, but albumin occurs in considerable quantity in the stools of typhoid and diarrhoea. In one instance much serum-albumin was found in the discharges of a chlorotic woman, who passed pale stools almost devoid of bile ; and once again in the case of a patient whose stools were deficient in bile, but who was not jaundiced (*v. Jaksch*).

3. Peptone.—For the detection of peptone, the fæces should be mixed with water so as to form a thin pulpy fluid, then boiled, and filtered while still hot. The filtrate will be clear or slightly tinged with red. When it has cooled, it may be tested for albumin with acetic acid and ferrocyanide of potassium. It usually happens that the liquid becomes a little turbid when the acetic acid is added (mucin), but the turbidity does not extend on the addition of ferrocyanide of potassium. When this is so, the mucin may be precipitated with acetate of lead, the filtrate tested in the manner afterwards to be described (chapter on *Urine*) with phosphotungstic acid, and the fluid which finally remains may be submitted to the biuret test. Should it happen that the fluid after boiling contains albumin as shown by the acetic acid and ferrocyanide of potassium test, this substance must be removed by combination with ferric acetate (see chapter on *Urine*), and the remainder of the process conducted as above.

The author has never found peptone in healthy fæces, but has met with it repeatedly in disease. He has collected the records of some fifty or sixty cases bearing upon this matter, and the results which were obtained depend upon seventy or eighty separate analyses.

Out of seven cases of typhoid, peptone occurred in the liquid stools in large quantities in five. Its presence remained doubtful in one, and it was absent from the discharges of the seventh.

It was present in all cases where the stools contained pus, as in dysentery (two cases), tubercular ulceration of the intestine (three cases), and suppurative peritonitis discharging pus into the intestine (one case).

In hepatic affections the character of the fæces in this respect was

very inconstant. In a series of cases of catarrhal jaundice with more or less acholic stools, no peptone was to be found ; whilst, on the other hand, the thin non-purulent discharges from a patient with syphilitic inflammation of the liver exhibited it in considerable quantity. It occurred plentifully in certain cases of atrophic cirrhosis and of carcinoma of the liver.

Acholic stools without jaundice were generally rich in peptone, but the results of analysis in these cases show much variety (*v. supra*).

4. Urea.—The presence of urea may be best ascertained by the method previously described (*vide Chapter I.*). For the estimation of metabolism, it is necessary to ascertain the total quantity of nitrogenous substances contained in the faeces. To do this, the stools should be treated with dilute acid (to prevent the evaporation of ammonia) and dried. The usual method for the analysis of organic compounds will then serve to determine the proportion of nitrogenous bodies present (see chapter on *Urine*).

5. Uric Acid and Xanthin Bases.—It would appear from *Weintraub's*¹⁸⁵ observations, that the faeces both of health and disease (in leukaemia) may contain uric acid and xanthin substances. (For their detection, see Chapter VII.)

6. Carbohydrates.—Various carbohydrates occur in the faeces. Of these, starch is the most prominent. It may be recognised at any time by the aid of the microscope. *Hoppe-Seyler*¹⁸⁶ asserts that grape-sugar and certain gummy carbohydrates may be contained in the stools.

To ascertain the presence of these substances, the faeces must be boiled with water, filtered, and the filtrate partially evaporated on a water-bath. One part of the fluid may then be tested with Trommer's or the phenyl-hydrazin test for sugar. To another part a little of the iodo-potassic-iodide solution may be added, to show the presence of starch. The remainder of the fluid may now be distilled, and the residue extracted with alcohol and æther (see chapter on *Urine*) ; the extract boiled with water and filtered ; the filtrate partially evaporated ; treated with dilute sulphuric acid, and boiled ; saturated with caustic soda ; treated again with cupric sulphate and boiled. The reduction phenomenon will show the presence of dextrin and gums (*Hoppe-Seyler*).¹⁸⁷ The property of benzoyl chloride to form insoluble compounds with carbohydrates in alkaline solutions may also be utilised as a test. It may be here mentioned that, in a series of hitherto unpublished observations made by the author on the occurrence of carbohydrates in faeces, no appreciable quantity of grape-sugar could be detected even in the most severe cases of diabetes, the result remaining negative, notwithstanding the exhibition of purgatives. (See the communication (which will appear shortly) by Rössler from the author's clinic.)

Of the pentose group, arabinose passes into the stools only when exhibited in large quantities, whereas xylose does so more readily, and rhamnose (methyl pentose) more easily than any other. The last-named was almost constantly detectable, in larger or smaller quantities, in the stools within a short time of its exhibition.

For the qualitative recognition of sugar the faintly acid fæces—acidified, if necessary, by the addition of sulphuric acid—is extracted by boiling with water, according to the methods detailed in Chapter VII. For quantitative estimations an aliquot portion of the fæces is extracted by boiling with a fifty-fold volume of water and filtered, the filtrate being then concentrated to 200 cc. and filtered again. The sugar present in this clear brown-coloured filtrate may be estimated by titration with Fehling's solution, in accordance with the regulations prescribed in Chapter VII.

7. Acids.

(a.) **Bile Acids.**—For the detection of bile acids, the distillation residue of the fæces may be tested in the manner already described in connection with the blood (p. 92). If biliary acids be present in great quantity, the application of Pettenkofer's test directly to a watery extract of the fæces (p. 92) will suffice for their recognition. They may also be detected by the addition of a watery solution of furfural and sulphuric acid (p. 92 and chapter on *Urine*).

(β.) **The Volatile Fatty Acids.**—To obtain these bodies from the fæces, the following plan may be adopted:—Dilute the faecal mass with water, add phosphoric acid, and distil. The distillate contains these acids together with indol, phenol, and scatol. If this distillate be neutralised with carbonate of soda and again distilled, the indol, phenol, and scatol will pass over, and the sodium salts of the fatty acids remain behind. Let these now be evaporated to dryness on the water-bath, the residue extracted with alcohol, and after the evaporation of the latter dissolved in water. The solution may then be tested for fatty acids.

The separation of the different fatty acids may be accomplished by fractional distillation when they are present in abundance. Moreover, they may be partially isolated by precipitation of the sodium salts with æther in alcoholic solutions of varying degrees of concentration (*v. Jaksch*).¹⁸⁸ When there is sufficient material at hand, it is a good plan to convert the acids into their silver or barium salts, and to effect their separation on the principle of their different degrees of solubility in water.

The estimation of the silver, barium, or sodium constituent in the corresponding salts may be effected in the reactions given below, and the proportion of the respective acids determined accordingly. Of the

volatile fatty acids butyric and acetic acids are those most readily to be found in the faeces.¹⁸⁹ Formic and propionic acids do not appear to have been recognised with absolute certainty, but we shall take account of them here, since it is certain, at any rate, that they occur in another secretion (urine). (See Chapter VII.)

(a.) **Formic Acid.**—This is a colourless liquid, of a pungent penetrating odour, which freezes at 0° C., boils at 100° C., and is miscible with alcohol and water.

1. Free formic acid is not precipitated by nitrate of silver, but this reagent will precipitate the alkaline salts of the acid from concentrated solutions. The silver compound blackens quickly in the cold, and when heated reduction takes place.

2. If a solution of perchloride of iron be added to a solution of a neutral salt of formic acid, a blood-red colour appears. This hue disappears on boiling, and a rust-coloured sediment remains.

3. If formic acid or an alkaline salt of the acid be heated with mercuric chloride to 60° or 70° C., a precipitate of subchloride of mercury forms. This reaction is impeded by the presence of free hydrochloric acid or of excess of an alkaline chloride.

(b.) **Acetic Acid** is a fluid with an acrid, pungent odour. Its boiling point is 119°, and it crystallises at 0° C. Heated with ferric chloride, its salts behave like those of formic acid. Nitrate of silver yields a precipitate in neutral solutions of a salt of acetic acid, and this precipitate dissolves in boiling water without being reduced.

When a salt of this acid is heated with sulphuric acid and alcohol, the characteristic odour of acetic æther is obtained.

(c.) **Propionic Acid.**—This is an oily fluid. It boils at 117° C. Propionic salts exhibit the same reactions with nitrate of silver as those of formic acid. With ferric chloride they do not yield a red colour.

(d.) **Butyric Acid.**—In the pure state butyric acid is an oily fluid, with a goat-like odour, which boils at 137° C. It is miscible in all proportions with alcohol and æther. Its salts, when treated with mineral acids, develop the characteristic goat-like smell. Ferric chloride solution does not give a red colour in solutions of such salts, while nitrate of silver forms a crystalline sediment which is insoluble in cold water.

To isolate butyric acid from the isobutyric acid in the faeces, the portion which distils at 158° C. should be treated with carbonate of guanidin (*Brieger*¹⁹⁰), and the guanidin salt obtained converted by heating into the corresponding guanamin. The base examined under the microscope will then exhibit the characteristic pointed rhomboids of the guanamin of isobutyric acid.

Valerianic, caproic, and others of the higher fatty acids are also present in the faeces. *Wegscheider*¹⁹¹ asserts that oleic, palmitic, stearic, capric, and caproic acids occur in the discharges of infants.

8. Phenol.—This substance is always a constituent of fæces. When the fatty acids have been converted into their sodium salts in the process described above for the separation of the volatile fatty acids, phenol passes over in the distillate. To isolate it from indol and scatol, the distillate must be rendered alkaline with caustic potash and again distilled. The phenol remains behind, and may be purified by distillation with sulphuric acid.

(1.) If a portion of the distillate now be treated with a solution of ferric chloride, a violet colour will show the presence of phenol.

(2.) The addition of bromine-water to another portion will cause the deposition of a crystalline sediment of tribromophenol.

(3.) Millon's reagent gives a red colour.¹⁹²

9. Indol and Scatol.—Both these bodies occur in the fæces, the latter having been detected there by *Brieger*.¹⁹³ To separate them from the phenol present, the distillate in the above process (see *ante*, *Volatile fatty acids*) should be treated with an alkali and again distilled, when indol and scatol will pass over. Indol forms in small colourless scales like those of benzoic acid. It dissolves in boiling water and very readily in alcohol, and is decomposed by strong alkalies.

Scatol, which also crystallises in colourless scales, is much less readily soluble in water, and possesses a disagreeable pungent smell. It is not decomposed by fairly strong alkalies.

To obtain either of these bodies separately, we avail ourselves of the lesser solubility of scatol in water, or of its property of resisting the action of alkalies.

(1.) If nitric acid which contains nitrous acid be added to a solution of indol, a distinct red colour is produced, or a red precipitate if the solution be concentrated.¹⁹⁴

(2.) Pine-shavings steeped in hydrochloric acid are turned red in an alcoholic solution of indol. The first test, when applied to scatol, does not produce a red coloration, but at most a slight turbidity. The second entirely fails. Jointly they will serve sufficiently for the discrimination of these two substances.¹⁹⁵

10. Cholesterin, Fats, and Non-Volatile Organic Acids.—

Cholesterin, as we have seen, seldom occurs as crystals in the fæces, but, according to *Hoppe-Seyler*, is in one form or the other an invariable constituent of those discharges. For its detection chemically, the residue which remains when the volatile fatty acids and phenol group have passed over in the process of distillation is to be treated with excess of sulphuric acid, and extracted first with alcohol and then with æther.

(1.) The æthereal extract is filtered, the æther removed by distillation, and the residue first digested with carbonate of soda on the water-bath, so as to remove any traces of the volatile fatty acids that may have

failed to pass over with the æther, evaporated to dryness, and again extracted with æther. (2.) The alcoholic extract is also filtered, treated with carbonate of soda, the alcohol distilled off, the residue dissolved in water, and finally, as before, extracted with æther. In the alkaline watery residue are contained the biliary acids (*v. supra*), oleic, palmitic, and stearic acids, which, according to *Hoppe*, may be separated by converting them into their barium salts.

Cholesterin and fat are taken up by the æther. The latter is evaporated, the residue treated with alcoholic solution of caustic potash, the alcohol removed by evaporation on the water-bath,¹⁹⁶ and the remaining fluid diluted with water and extracted with æther. The fats remain as soaps in the watery solution, while the cholesterin is dissolved by the æther.

Cholesterin may be recognised by the following tests :—

(1.) A little concentrated sulphuric acid applied to the crystals on a slide will cause them gradually to colour a reddish yellow round their borders, and finally to grow smaller and disappear.

(2.) When the crystals are dissolved in chloroform and sulphuric acid is added, a blood-red colour forms, presently changing to purple-red. The sulphuric acid at the same time shows a strong green fluorescence.

(3.) A particle of cholesterin to which a little nitric acid has been added is placed in a small dish and evaporated to dryness in the water bath. It leaves a yellow stain, which turns a yellowish red on the addition of ammonia (*Schulze*).¹⁹⁷

The soap-solution obtained in the above process is to be rendered acid with dilute sulphuric acid, and the resulting fatty acids removed by filtration. If the filtrate be neutralised with ammonia, evaporated, and extracted with alcohol, it will be found to contain glycerine. The fatty acids may be again dissolved in æther, repeatedly saponified,¹⁹⁸ collected and dried, and then identified, first by ascertaining their melting-points, and again by converting them into their barium salts, and determining in each case the value of the barium constituent (*Hoppe-Seyler*).¹⁹⁹

Fats, and especially neutral fats (tri-glyceride), soaps, non-volatile fatty acids, and cholesterin are present in all faeces. Acholic stools present them in relatively large quantities. According to *Müller*,²⁰⁰ clinical inferences can be based on a knowledge of the melting and solidifying points of the fatty acids.²⁰¹ Both points are higher in proportion to the efficiency of intestinal absorption. When the fatty acids of the faeces contain a proportion with a melting-point so low as 50° C., this function is probably impaired.

For the quantitative estimation of fats in the stools the methods of *Hoppe-Seyler* and *Benedikt*²⁰² may be employed. That of *Müller*²⁰³ is

more expeditious, and serves well for clinical purposes. To determine the amount of fats formed with a given diet, as milk or flesh, powdered animal charcoal is administered with the first meal, which should be taken on an empty stomach. The earliest resulting fæces are thus stained black. A portion is dried at 100° C., and by extraction with ether in *Sorobet's* apparatus the neutral fats and free fatty acids are obtained. These are dissolved in warm alcohol and a little ether, and submitted to titration with phenol-phthalein solution and alcoholic solution of caustic potash. The result of titration shows the proportion of fatty acids present. Another portion of the dried fæces is extracted first with acid alcohol and afterwards with ether to determine the amount of soaps contained in it. In intestinal disease, especially such as involves the lymphatics, and where the flow of bile is suspended (*Muller*),²⁰⁴ the fæces contain great quantities of fat, showing that absorption is impeded.

11. Colouring Matters.

1. Urobilin.—This is the normal colouring matter of the fæces (p. 194). It can readily be obtained from the stools by treating them with acid alcohol.

*Mehu's*²⁰⁵ method for the isolation of urobilin in fæces yields excellent results. A watery extract of the fæces is made, and to this is added sulphuric acid in the proportion of 2 grms. to the litre, and solid ammonium sulphate. The fluid is then filtered, and the precipitate washed with a warm saturated solution of ammonium sulphate, dried on a water-bath, and extracted with a boiling alcoholic solution of ammonia. *Ad. Schmidt*²⁰⁶ recommends for the detection of urobilin (hydro-bilirubin) that the fæces be treated with a strong aqueous solution of corrosive sublimate; if present, a red colour is soon developed. Urobilin may be identified by its characteristic spectrum in acid solutions. This exhibits a well-defined absorption-band between the lines *b* and *F* (*Fraunhofer*) of the solar spectrum (fig. 137).

It should be noted that urobilin may be present even in acholic stools.

Urobilin disappears from the stools in phosphorus poisoning (*Riva*,²⁰⁷ *Lanz*²⁰⁸). Its reappearance there is a favourable symptom.

2. Blood Colouring Matter.—Pure blood occurs in the stools only after very profuse and rapid haemorrhage into the intestine. In all other cases where its constituents are found they are greatly altered. Haematoxin crystals are rarely seen. Haematin is the form in which blood pigment occurs most commonly. This substance may best be recognised by Teichmann's test, or with the spectroscope (p. 73).

3. Bile Pigment.—The fæces never contain bile pigment in health, but it is found abundantly in the discharges in cases of catarrh of the

small intestine. It may be best detected by the application of *Gmelin's* test (nitric acid). On the addition of a little impure nitric acid to a specimen of faeces in which bile pigment is present, the mass changes colour quickly, and the separate drops of the acid are surrounded with rings of green, red, and violet. The appearance of a green ring is very characteristic of bile pigment, and is due to the formation of biliverdin. We have already spoken of the other pigments which may occur in the stools (see p. 193).

12. Intestinal Gases.—These consist of hydrogen, carbonic acid, nitrogen, and volatile carburetted hydrogen (methan).²⁰⁹ It is not yet definitely determined whether sulphuretted hydrogen is formed in the intestine or not. *Senator* and *Ottavio Stefano*,²¹⁰ however, maintain that in certain morbid states this gas is generated in such quantity as to cause symptoms of poisoning. The fact that sulphide of bismuth is formed in the alimentary canal when nitrate of bismuth has been taken (*v. Jaksch*), lends probability to the assumption that sulphuretted hydrogen is also produced there. According to *Hammarsten*,²¹¹ the latter occurs in small quantity in normal faeces. *Nencki*²¹² has detected methylmercaptan in the faeces.

13. Ptomaines.—Putrescin and cadaverin are present in the stools, and may be recognised by the methods already laid down. Moreover, the ptomaines known to be produced by pure cultivations of certain pathogenic fungi, have recently been obtained directly from the faeces, as by *Pouchet*²¹³ in cases of cholera. *Baumann* and *Udransky*, *Stadthagen* and *Brieger*,²¹⁴ have separated diamines from the stools of patients with cystinuria (see chapter on *Urine*). They believe that these substances are absent from normal faeces. See the methods described at p. 187.

14. Ferments.

Diastase and **Invertin** are generally present in the stools of healthy children (*v. Jaksch*).²¹⁵ For the detection of diastase see Chapter II.

B. Inorganic Substances.—The consideration of such inorganic substances as assume a crystalline form has already engaged our attention (see *Crystals*). Chloride of sodium may be detected in the faeces thus:—An extract is made with water, acidulated with nitric acid, filtered, and the filtrate tested with nitrate of silver. A white precipitate (chloride of silver), soluble in ammonia, will show the presence of the sodium salt.

In *Hoppe-Seyler's*²¹⁶ method for the quantitative analysis of inorganic matter in the stools, the substances which are soluble in alcohol are separated from those which are soluble in dilute acetic and hydrochloric acids before the process of incineration is commenced. If this

is not done, there is danger that the nuclein, which is nearly always present in fæces, will be decomposed, setting free its phosphoric acid, which may then either remain uncombined or displace other acids from their compounds. The analysis, both quantitative and qualitative, of the incinerated ash is conducted according to methods which are sufficiently familiar.²¹⁷

IV. EXAMINATION OF THE MECONIUM.—The term "meconium" is applied to the substance discharged from the rectum of the child immediately after birth. It is a thick, sticky, viscous fluid, of a greenish-brown colour. When examined by the microscope, meconium exhibits some intestinal epithelium cells, fatty particles, both fluid and solid,²¹⁸ numerous cholesterin crystals, a quantity of more or less well-formed crystals of bilirubin, and some downy hairs. There are immediately after birth no fungi, and (according to *Escherich*²¹⁹) no spores. After the lapse of twenty-four hours, however, the discharges exhibit a very different character. They now contain abundance of micro-organisms, and Escherich obtained from them by Koch's plate-cultivation methods three distinct microbes.

After the child has taken the breast, the bacteria of the stools, according to the same authority, are represented by two species of micro-organisms. The first consists of thick, curved, rod-like bodies, measuring 1-5 μ in length by 0.3-0.4 μ in thickness. The other is a micro-organism which closely resembles the lactic acid bacillus of *Hüppe*.²²⁰ *Baginsky* has obtained similar results.

In addition to the above, the meconium contains numerous squamous epithelium cells, derived from the pharynx and œsophagus, or from the anal orifice (*Bizzozero*).²²¹

Ziceifel and *Hoppe-Seyler*,²²² who have investigated the chemical constitution of the meconium, found it to contain bilirubin, biliverdin, and biliary acids, but no hydro-bilirubin (urobilin). *Wegscheider*²²³ found traces of peptone, fats, and soaps, bilirubin, and traces of hydro-bilirubin in infantile stools. In a specimen which the author examined, there was no serum-albumin, peptone, or sugar. There was abundance of mucin. Bilirubin was the only pigment present.

V. CHARACTER OF THE FÆCES IN CERTAIN INTESTINAL AFFECTIONS.

1. Acute Intestinal Catarrh.—In this condition the quantity of the stools is subject to great variety. They are usually fluid, thin, and slimy, yellowish brown in colour, and emit a most unpleasant smell. Their reaction is alkaline, except in the case of acute enteritis of children, when it may be acid. Such stools usually contain great quantities

of mucus, and there are often visible to the naked eye food remnants in great quantity.

Microscopical examination reveals an abundance of fungi of various descriptions, large quantities of intestinal epithelium, and isolated leucocytes.²²⁴

2. Chronic Intestinal Catarrh.—In this disease the stools exhibit no very distinctive characters, whether to the naked eye or microscopically.

*Nothnagel*²²⁵ lays down the following rules for the localisation of chronic idiopathic intestinal catarrh, according to the character of the faeces :—

1. When the large intestine is alone involved, a single discharge takes place within twenty-four hours. Diarrhoea, however, is apt to recur at certain regular intervals.

2. When the small intestine alone is engaged, the motions are also likely to be sluggish.

3. When both the large and small intestines are the seat of catarrh, continuous diarrhoea is apt to ensue.

4. Solid or semi-solid stools containing hyaline particles of mucus, which can be recognised only with the microscope (see p. 195), and devoid of mucus visible to the naked eye, point to implication of the upper part of the large intestine.

5. The presence of bile pigment in the stools, as shown by Gmelin's test, invariably indicates a catarrh of the ileum and jejunum. In such cases also the faeces are usually found to contain epithelial cells and mucus, deeply stained yellow by the bile-colouring matter.

In certain forms of chronic catarrh, where the large intestine is especially involved, it sometimes happens that the bodies described at p. 195 are to be found in the stools. Such an affection is then called enteritis tubulosa or membranacea ; but it is probable that these manifestations accompany other sufficiently dissimilar morbid states. Our present knowledge of the subject is defective.

3. Ulcerative Enteritis.—The diagnosis of this condition is always attended with difficulty. It is usually (though not always) accompanied with diarrhoea. In a questionable case, the appearance of blood in the stools makes ulceration probable ; but we cannot derive any conclusive evidence from either the physical or the chemical character of the faeces to establish the diagnosis. A diligent examination of the discharges, however, may in certain specific forms of ulceration disclose the presence of the pathogenic organisms to whose influence the process is known to be due. The detection of the tubercle-bacillus especially, is in this way a fact of the utmost clinical significance (see p. 213).

4. Typhoid Fever.—This disease is usually characterised by abundant foul-smelling discharges of the colour of pea-soup. They contain large quantities of bile pigment, a fact which points to a catarrh of the small intestine, and to which also Nothnagel attributes the peculiarly offensive character of the smell emitted.

The reaction of typhoid stools is in all cases alkaline.

Microscopical examination shows numbers of bile-stained epithelial cells, some leucocytes, abundance of triple phosphate crystals, and a profusion of fungi. Nothnagel's clostridia are especially prominent amongst these. The typhoid-bacillus, of course, infests the discharges of this disease; but it cannot be distinguished from the other micro-organisms by a simple microscopical examination. This can be done only by the bacteriological methods indicated in the present chapter.

The stools of typhoid in its later stages may be those of intestinal ulceration. Thus, when haemorrhage results from the extension of typhoid ulcers, the fæces will be blackened, and yield the chemical reactions which denote the presence of a derivative of blood pigment (haematin).

5. Dysentery.—The discharges of dysentery are subject to a great variety of character; but there is one respect in which they are constant, for they always contain abundance of mucin, and in the author's experience also some serum-albumin and much peptone.

Under the microscope there are to be seen great quantities of leucocytes, intestinal epithelium, and fungi. Tolerably perfect red blood-corpuscles are occasionally visible. The number of these latter varies within broad limits, but the other microscopical appearances are remarkably uniform.

The grosser properties of dysenteric stools, on the other hand, display notable differences. Founded upon these, *Heubner*²²⁶ distinguishes:—

1. *Mucous and muco-sanguineous discharge.*—A pale yellow, viscous, transparent substance tinged with blood, cohering in masses, with or without admixture of fæces.

2. *Sanguineo-purulent discharge.*—A reddish or yellow fluid, containing flocculent or solid particles as large as a pea or a bean. Such stools may be compared to raw minced meat.

3. *Discharge of pure blood.*—This occurs when a vessel has been opened by the extension of a dysenteric ulcer.

4. *Discharge of pure pus.*—This consists almost exclusively of leucocytes, and belongs to the later stages of dysentery.

5. *Gangrenous stools.*—Such stools are brownish-red or brown-black in colour, from the presence of altered pigment. They emit a putrid odour. They indicate extensive gangrene of the intestinal mucous membrane.

It was in dysentery that the stools were first noticed to contain those mucous particles which have been compared to frog-spawn (see *Naked-eye Characters of the Faeces*), and which *Nothnagel* afterwards observed in other intestinal diseases. They have no special clinical significance in this disease.

On the whole, it may be said that the naked-eye characters of dysenteric stools are so remarkable that they will ordinarily suffice to establish a diagnosis without the aid of the microscope.

Amœbæ, which have lately been found in such stools, have been credited with a causal relation to the disease (*Hlava*, *Kartulis*, *Korazz*, and *Viraldi*).²²⁷ A similar importance is attached by others to a pathogenic fission-fungus (*Klebs*, *Chantemesse*, *Widal*, and *Arnaud*),²²⁸ but with less reason. The occurrence of amœbæ in dysenteric stools has been observed in so many quarters that it is probable there is some connection between the organism and the disease. Still, amœbæ are also found in health (*Schuberg*),²²⁹ and it is possible that various micro-organisms may be concerned in exciting the symptoms of dysentery. *O. Arnaud* and *Lareran*²³⁰ attach importance to the *Bacterium coli commune*. *De Silvestri*²³¹ obtained organisms from the dysenteric stools during an epidemic, and ascertained that they were pathogenic in animals.

6. Cholera.—During an epidemic of cholera there is usually prevalent a form of diarrhoea which is distinct from that disease, and it is of the utmost importance to possess the means of discriminating between the two. The discharges of the less formidable complaint are not characterised by any special changes; but, in a doubtful case, the investigation in the stools for cholera-bacillus (as indicated in this chapter) may be needed to establish the diagnosis.

In a pronounced case of Asiatic cholera, on the other hand, no kind of ambiguity can exist. The discharges are thin, and devoid of smell and colour. They have been aptly termed "rice-water" stools. Microscopically they abound in leucocytes and epithelium, and their specific micro-organism, the comma-bacillus, may be readily detected. It must, however, be borne in mind that "rice-water" stools are not by themselves pathognomonic of cholera. They are seen repeatedly in heat-apoplexy and arsenical poisoning; and in such connection, as well as in cholera, they hold a profusion of intestinal epithelium. It follows that the diagnosis of Asiatic cholera will rest on an absolutely secure basis only when the comma-bacillus has been found, separated from the stools, and cultivated by the methods with which we are already familiar (p. 205). Chemically, the discharges of cholera contain serum albumin²³² and much mucus.

7. Hæmorrhagic Stools.—Blood is discharged with the stools in cases of great venous congestion of the intestine, in typhoid, in tubercular

and dysenteric ulceration of the stomach or intestine, and in round ulcer of the stomach or duodenum. These cases are always attended with symptoms of severe intestinal trouble. The blood is usually profoundly altered (see this chapter, *Red Blood-cells*) ; but when the hæmorrhage has taken place in the lower part of the alimentary canal, as the sigmoid flexure or rectum, pure bright blood may be passed.

8. Acholic Stools.—The stools may be deficient in bile in cases of jaundice from obstruction of the biliary ducts, or they may be so in the absence of this condition.

They are characterised by (1) their whitish-grey colour, (2) the abundance of fat which they contain, and (3) a profusion of fat crystals, probably soaps of soda, lime, and magnesia (fig. 96).

Such stools in connection with jaundice imply an obstruction to the flow of bile by blocking of the ducts. When they occur independently of obstruction, the underlying cause is not yet sufficiently understood. Many theories have been framed to account for the phenomenon :—(1) It may be either that the bile pigment has undergone some change in the intestine which prevents the formation of its metabolic product (urobilin) ; or (2) the secretion of bile may be so scanty that there is not enough pigment for the elaboration of urobilin ; or (3) it is possible in such cases that the latter is replaced by certain colourless metabolites or chromogens of bilirubin (*v. Nencki's leuco-urobilin*). The latter view is supported by the fact that considerable quantities of urobilin may be obtained from acholic stools by extraction with acid alcohol (*v. Jaksch, Pel, Le Nobel*).²³³ The stools may be devoid of bile in cases of the most varied origin—as in tuberculosis of the intestine, chronic nephritis, and chlorosis—where no trace of jaundice is present. They are commonly so in the fatty discharges which accompany indigestion in children (*Biedert*).²³⁴ *Berggrün* and *Katz*²³⁵ have observed acholic stools in the chronic tubercular peritonitis of children. In these cases, as usual, the fæces contained excess of fat. It follows, therefore, that we cannot infer the character of the stools from the presence or absence of this symptom. But in all cases where colourless stools concur with jaundice, the cause is to be found, as has already been said, in obstruction of the biliary passages.

CHAPTER VII

EXAMINATION OF THE URINE

THE urine is the secretion of the kidneys.¹* A sufficient and exhaustive knowledge of the characters of this secretion is a point of the utmost consequence to the physician, since the changes which it undergoes are the expression of numerous morbid processes, and their intelligent interpretation affords the surest aid to diagnosis.

I. NAKED-EYE INSPECTION OF THE URINE.

1. Quantity.—The quantity of urine secreted in health varies within broad limits, and depends at any time upon the relation subsisting between the imbibition and abstraction of fluids in the system. It follows that an error as to excess or deficiency can be considered morbid only when very marked. In general terms it may be said that a healthy able-bodied man will pass 1500 to 2000 cc. of urine in twenty-four hours. The rate of secretion varies at different periods of the day. In the early hours of the night the urine is abundant and of relatively low specific gravity; later it is scantier and more concentrated, while during the waking hours it again becomes more abundant (*Wollheim de Fonseca*).² Secretion is diminished in sleep (*Glum*).³

Under pathological conditions the healthy standard may be widely departed from in either direction.

In order to estimate the quantity of urine secreted, that which is passed within twenty-four hours should be collected; and it is well to date the period from eight o'clock of one morning to eight o'clock of the next. When only half this interval is taken—and especially when the estimation of urea is in question—it is important that the bladder should be previously emptied. The patient should be admonished to make water before going to stool, and even then a certain allowance must be made for urine passed in the act of defæcation. Should the patient be the subject of incontinence, it is very difficult to form an

* We shall confine ourselves here to a description of the simpler processes, and such as are likely to be applied clinically. For more exhaustive information the reader is referred to the text-books of urinary chemistry (see references 1 and 4).

accurate estimate of the quantity of urine secreted; and this can be done only by passing the catheter as often as possible—hourly, if it may be—so as to minimise the escape of fluid. In paralysis of the bladder, while sensation remains unimpaired, the escape of urine may be prevented by the use of a permanent receptacle. The urine saved should, in any case, be placed in a vessel of two litres capacity, and provided with a graduated scale showing its capacity in cubic centimetres. Its quantity may be most accurately estimated by weighing it.*

A diminution of the quantity of the urine (oliguria) occurs in febrile conditions, in disturbances of the circulation of all kinds, and especially of the capillary circulation, in acute, and in some forms of chronic nephritis. The urine is increased in quantity in diabetes mellitus, diabetes insipidus, contracted kidney, amyloid degeneration of the kidney, and usually in convalescence after acute diseases. [To these causes may be added rare cerebellar disease, hysteria, and nervous conditions.] Under the heading of acute diseases the increase is most pronounced in the non-febrile period of relapsing fever, at the termination of an attack of acute nephritis—whether cure be impending or the acute is passing into the chronic form of the disease—and in the restoration of the balance in the capillary circulation, as where compensatory changes take place in heart-disease, &c. Finally, the renal secretion is promoted by certain drugs, as the salts of acetic and salicylic acid, digitalis, calomel, and diuretin (sodio-salicylate of caffein).

A complete suppression of the urinary secretion (anuria) is an accompaniment of uræmia and of all diseases that are attended with the abstraction of water from the system: such are severe acute anaemia, gastric and intestinal catarrh, cholera, and dysentery, and it occurs also in certain toxic states, as poisoning by oxalic acid and arsenic. [A remarkable case⁵ has been recorded in which total anuria lasting for a week was attributed to the impaction of a calculus in one ureter, causing obstruction of that, and suppression by reflex influence; there were slight symptoms of uræmia shortly before recovery.] The transitory suppression of urine which sometimes occurs in healthy persons after profuse perspiration, and lasts only a few hours, is altogether physiological. A simple increase or diminution of urine will not by itself establish the nature of a disease, but it is always an important factor in diagnosis, and we shall see by-and-by that the consideration of this point will enable us especially to discriminate between certain forms of kidney-affection.⁶

2. Specific Gravity (Density) of the Urine.—The density of the urine varies greatly in health, and is for the most part in inverse ratio to its quantity. If we assume the latter to be on an average

* See the text-books referred to, reference 4.

1500–2000 cc., then the sp. gr. of healthy urine may be stated at 1.017 to 1.020. It may be estimated most accurately by means of the pycnometer (see text-books),⁷ but for clinical and practical purposes an instrument constructed on the principle of the hydrometer will serve well. Such an instrument, when employed for testing urine, is called a urinometer. It is well to have two in use, one for taking the specific gravity when this lies between 1.000–1.025, the other where it is as high as 1.025–1.050.

A serviceable urinometer should be so made that the degrees on the index-scale shall be separated by a sufficient interval not less than 1 mm.; and when very accurate results are aimed at, it should be graduated in tenths. Moreover, it should be furnished with a thermometer, also graduated in tenths of a degree, and recording temperatures between 0° and 30° C.

A new urinometer should be tested in distilled water. If accurate, it will sink to the mark 1.000, where the scale is graduated so low as this.

The specific gravity is tested in the following manner:—The urine is poured into a cylindrical glass vessel of suitable width. Should froth form, it must be removed either with filter-paper, or by filling the vessel to the brim, when it may be blown off. The urinometer is then placed in the urine, and care is taken that it is not anywhere in contact with the sides of the vessel. When it is quite stationary in the liquid, the specific gravity is read off, the observer bringing his eye on a level with the concave surface of the liquid. The mark on the scale which corresponds to the lowest point of this concavity will indicate the specific gravity of the fluid.

Greater accuracy may be attained by testing the urine at some particular temperature, for which the urinometer has been constructed.

[According to *Thudichum*,⁸ the average of a series of analyses gave the specific gravity of healthy urine as 1.020, the quantity passed in the twenty-four hours appearing in the same series 1400–1600 cc. or 48–56 fluid ounces.]

An abnormal specific gravity of the urine is a fact of great importance in disease. It affords an approximate estimate of the quantity of solids excreted by the kidneys, and consequently of the energy of the metabolic processes within the system. It may be stated as a general rule, that when the quantity of the urine is diminished in disease, its specific gravity is raised. A considerable departure from this rule implies one of two things:—Either tissue-changes are notably suspended, and their products, urea, uric acid, &c., formed in smaller quantities; or these processes remaining active, such products fail to be removed by the kidneys. To the first of these causes is to be assigned that rapid

decline in the density of the urine which sometimes precedes a fatal termination in acute fevers. Of still more serious import is a sudden fall in the specific gravity in nephritis, unattended with any alteration in the quantity of urine passed. The phenomenon in this instance points to the failure of the diseased kidneys to separate the urea and salts elaborated within the system. The author has had many opportunities of observing that such a fall in the specific gravity is apt to precede—usually by several days—the oliguria and suppression which herald an attack of uræmia; and it often affords a valuable warning of what is impending at a time when all other symptoms are wanting. Moreover, the symptoms of uræmia may develop whilst the urine remains but little diminished in quantity, and in such cases we shall always find that its specific gravity is greatly lessened.⁹

3. The Colour of the Urine.—The normal urinary pigments have not yet been separated. *C. Vierordt*¹⁰ concludes from spectroscopic appearances that they are several in number. [Attempts to separate the colouring matter of normal urine were made by *Tichborne*¹¹ and *Thudichum*.¹² The latter gave to the substance derived from urine by his process the name of urochrome. *Garrod*¹³ has lately, by a new process, isolated from the urine a substance to which he believes the colour is entirely or almost entirely due. He believes this to be a definite chemical body, and distinct from the urinary products, such as urobilin and hæmatoporphyrin, which possess spectroscopic properties.] Up to the present, on the other hand, but two chromogens have been determined in the urine, viz., indican (sulphate of indoxyl; see *Indicanuria*), and the chromogen of urobilin.¹⁴

[*Garrod*¹⁵ has shown that hæmatoporphyrin as well as urobilin is present in healthy urine; but both are in such small quantities that they cannot be held to impart its colour to the fluid.

Four pigments are usually distinguished as belonging to the urine (*F. Taylor*). These are:—1. Normal urobilin. 2. Febrile or pathological urobilin, which is identical with stercobilin (*MacMunn*). 3. Urohæmatoporphyrin, derived from hæmin. 4. Uroërythrin, which is the pigment of pink urates.

The chromogens of the urine are substances which develop a colour on the addition to the fluid of some oxidising or other reagent. These are likewise four, namely:—The chromogens (1) of indigo (indican); (2) of pathological urobilin; (3) of anaemia; (4) of melanin. *MacMunn* believes further that normal urobilin, which is the principal pigment of healthy urine, exists there in part as a chromogen.]

The colour of healthy urine depends upon its degree of concentration, being darker as this is more pronounced.

The same statement holds in general for disease, but with notable

exceptions; for there are some affections in which a high colour and an abundant flow coincide, and others in which the urine is at once pale and scanty.

In some diseases, and especially in fever, additional colouring matters are secreted. The nature of some of these is a matter of speculation (*uroërythrin*, *urochrome*).¹⁶

At some period in the course of a disease, the urine may undergo a change of colour from the admixture of blood. When the latter is present in small quantity, the secretion may be flesh-water-coloured. When blood is more plentifully effused, it may be a bright ruby-red (see *Hæmaturia*).

Bile pigment imparts a brownish-yellow or green tint. Its presence may usually be detected by shaking up the urine, when a yellow foam will form upon it. It must, however, be borne in mind that urine which contains much *urobilin* will yield a similar foam (*Leo Liebermann*¹⁷); in the latter case the secretion is always of a dark brown-red (see *Urobilinuria*). An excess of indoxyl sulphates will cause the urine to assume a dark-brown hue, and the yellow foam will not form upon it (see *Indicanuria*). The brown tint in such cases is due, not to the indoxyl salts, which are colourless, but to some other substance which is present with them. Urine rich in urobilin always has an intensely brownish-red colour (see *Urobilinuria*).

Certain drugs, too, will affect the colour of the urine. Rhubarb and senna cause it to become brown or blood-red; a black colour is developed when carbolic acid is taken into the system, especially if the urine be allowed to stand for some time, and the same appearance follows the exhibition of naphthalin, hydrochinon, resorcin, and pyrocatechin. The blackening of the urine by carbolic acid is ascribed by *Baumann* and *Preusse*¹⁸ to the formation of the oxidation-products of hydrochinon. The use of quinine, kairin, antipyrin, and thallin, and sometimes sulphonal (see *Hæmatoporphyrinuria*), also colours the urine more or less deeply.

It may be stated in general that the urine is darkly coloured in fevers, and in congestion of the kidneys due to heart-disease, emphysema, &c. It is, on the other hand, deficient in colouring matter in diabetes mellitus and insipidus, chronic nephritis, urina spastica, and all kinds of anaemia. [The urine of pernicious anaemia, though of low specific gravity, not exceeding 1016 (*Hunter*), is usually of very high colour (*Fayje*,¹⁹ *Mott*,²⁰ *Hunter*²¹). The urine of phthisis has a tendency to become dark on standing, and sometimes turns quite black (*Hale-White*).²²] In cancer, especially when it implicates the intestinal canal, the urine is apt to be dark and pigmented. In such cases an excess of indican may generally be determined.

Vogel has endeavoured to construct a standard scale for the estimation of colour in the *urine*.

[The following table from *Halliburton*²³ shows the nature and origin of the chief variations in tint :*—

Colour.	Cause of Colour.	Pathological Condition.
Nearly colourless.	Dilution or diminution of normal pigments.	Various nervous conditions, hydruria, diabetes insipidus, granular kidney.
Dark-yellow to brown-red.	Increase of normal or occurrence of pathological pigments.	Acute febrile diseases.
Milky.	Fat globules. Pus corpuscles.	Chyluria. Purulent disease in urinary tract.
Orange.	Excreted drugs, e.g.	Santonin, chrysophanic acid.
Red or reddish.	Unchanged haemoglobin. Pigments in food (logwood, madder, bilberries, fuchsin).	Hæmorrhage or hæmoglobinauria.
Brown to brown-black.	Hæmatin. Methæmoglobin. Melanin. Hydrochinon and catechol.	Small hæmorrhages. Methæmoglobinuria. Melanotic sarcoma. Carbolic acid poisoning.
Greenish-yellow, greenish-brown, approaching black.	Bile pigments.	Jaundice.
Dirty green or blue.	A dark blue scum on surface with a blue deposit, due to excess of indigo-forming substances.	Cholera, typhus ; seen especially when the urine is putrefying.]
Brown-yellow to reddish brown, becomes blood-red on addition of alkalies.	Substances introduced into the organism with senna, rhubarb, and chelidonium.	

4. The Reaction of the Urine.—Healthy human urine is ordinarily acid. The reaction is due not to free acid, but to the acid salts (phosphates and urates) which it contains.

It is, however, subject to considerable variations in this respect. *Quincke*²⁴ has determined that the acidity is in general less in the fore-

* The author is of opinion that neither *Vogel's* table nor that of *Halliburton* is of the slightest value ; and in fact considers that they are likely to lead to superficiality in the examination of this important secretion. Nevertheless, in deference to the views held by his deceased friend *Cagney*, he has retained this table in the present English edition, though he has not adopted it in the German edition.

noon than at any other time, and the urine of healthy persons may even exhibit an alkaline reaction then.²⁵

The reaction of the urine is also modified by diet. It may be alkaline after an ample meal, or the ingestion of alkalies and of substances such as the salts of acetic, tartaric, and citric acids, which are converted into carbonates in the system. The administration of acids, on the other hand, renders the acidity of the urine more marked.

Healthy urine, when allowed to stand for some time, becomes [first more acid from an increase of acid phosphates, as well as lactic and acetic acids, and then] alkaline in consequence of the action of a micro-organism, the *micrococcus ureæ* (see p. 273), which decomposes the contained urea into carbonate of ammonia. [The urine of phthisis remains acid for a very long time, sometimes even for four months (*Hale-White*).]²⁶ It occasionally happens that the same urine will turn red litmus paper blue, and blue litmus red, i.e., it is amphoteric. This depends upon the presence in it of acid or neutral phosphates (*Huppert*).²⁷

In different morbid states the freshly-passed urine may be either alkaline or acid ; but its reaction in disease is valuable as a symptom only when the causes which are known to influence it in health can be excluded. A fact of the greatest consequence in this connection is the alkalinity due to ammoniacal fermentation of urea or of uric acid,²⁸ and this can usually be ascertained directly by the sense of smell. The urine is, as a rule, acid in febrile conditions, diabetes, and leukæmia ; in scurvy, too, it is apt to be intensely acid. On the other hand, it is alkaline in simple and pernicious anaemia and chlorosis. According to *Bence Jones*, the alkalinity in these cases depends upon the deficient formation of acid in the stomach. In chlorosis the point is of interest to the physician, insomuch that whilst the reaction of the urine continues to be alkaline, it may be inferred that the morbid process on which it depends is still going on. Ammoniacal urine implies ammoniacal fermentation within the bladder. This may be due to the use of an unclean catheter, and commonly arises in the course of cystitis. [Alkalinity due to the presence of ammonia may be distinguished from that caused by fixed alkalies from the fact that litmus paper turned blue by it again becomes red when dried in a gentle heat.]

The reaction of the urine is best tested with red and blue litmus paper. The comparative estimation of acidity may be effected by *Huppert's* method.²⁹ [The acidity of the urine during twenty-four hours is equivalent to about 14 grs. of carbonate of sodium or to 30 grs. of oxalic acid (*Taylor*).³⁰]

*D. Turner*³¹ proposes to apply *Kohlrausch's* telephone test for determining the electrical resistance of the urine, as a resource in diagnosis. The greater the resistance offered to the current by the urine, the more satisfactory it would seem is the health of the person under observation.

5. Odour.—Fresh healthy urine has a peculiar smell, somewhat resembling that of hay. The urine of diabetics is faintly aromatic ; that loaded with acetone has a fruity odour. Decomposition is attended by the formation of ammonia, which is easily recognised. The administration of turpentine or myrtol gives the secretion the scent of violets. The unpleasant smell of the urine after the ingestion of asparagus is due, according to *Nencki*,³² to the presence of methylmercaptan.

II. MICROSCOPICAL EXAMINATION OF THE URINE.—

Healthy urine is generally quite clear when first passed. On standing, if not decomposed by the rapid development of fungi, it deposits a filmy cloud. Microscopically this deposit is seen to consist of a few crystals of various kinds, some white blood-corpuscles, and epithelial débris.³³ But the appearance of freshly passed urine may differ greatly from this, even in health. The concentrated urine passed in the morning sometimes deposits an abundant sediment of urates altogether independently of disease.

The examination of morbid urine affords information of the utmost consequence in diagnosis. Such urine may either be turbid when passed, or it may deposit a variable quantity of sediment when allowed to stand for some time ; and when this sediment is looked at through the microscope, it is seen to hold certain substances of very variable character. For convenience of description we shall divide these substances into two classes, the *Organised* and the *Unorganised* urinary deposits.

The microscopical examination of the urinary sediment may be conducted in the following manner :—The urine is allowed to settle, the clear supernatant fluid poured off, and some of the sediment placed in a conical glass, when it is again allowed to stand for a while. A little is then removed with a pipette and placed on a slide for examination. If the sediment should be scanty and require a long time—say twenty-four hours—to deposit, it should be set apart in a cool place, so as to check the excessive development of fungi and fermentative processes, which might alter the character of the specimen. It is well also to add some indifferent antiseptic substance to the urine, as thymol, hydriodic acid, or oil of turpentine. An admirable admixture is that of *Salkowski* :³⁴ 20 to 30 cc. of a fluid containing 5 to 7.5 cc. of chloroform in a litre of water. Carbolic acid should not be employed, because it will cause a precipitate with any albumin which may be present.

The process may be rendered more certain and expeditious by the use of *Stenbeck's* sedimentator (*v. Jaksch, Litten*³⁵), the nature of which is explained by the accompanying figure (fig. 98). As used by the author, it is fitted with a treadle-wheel, and worked by the foot instead of the hand. It is also protected by a wooden case, within which the centrifugal apparatus is made to rotate. This is a precaution against accidents. With this instrument a few minutes is sufficient to produce a deposit even in urine which contains but little sediment, and where it is avail-

able no other procedure is required. The sediment when deposited is withdrawn by means of a pipette and submitted to microscopical examination as already directed. After some years' experience of an instrument of this kind, the author can speak in the highest terms of its usefulness, in spite of *Allin's*²⁶ depreciatory comments. *Fr. Winkler* and *J. Fischer*²⁷ have employed a galvanic current to precipitate the urinary sediment.

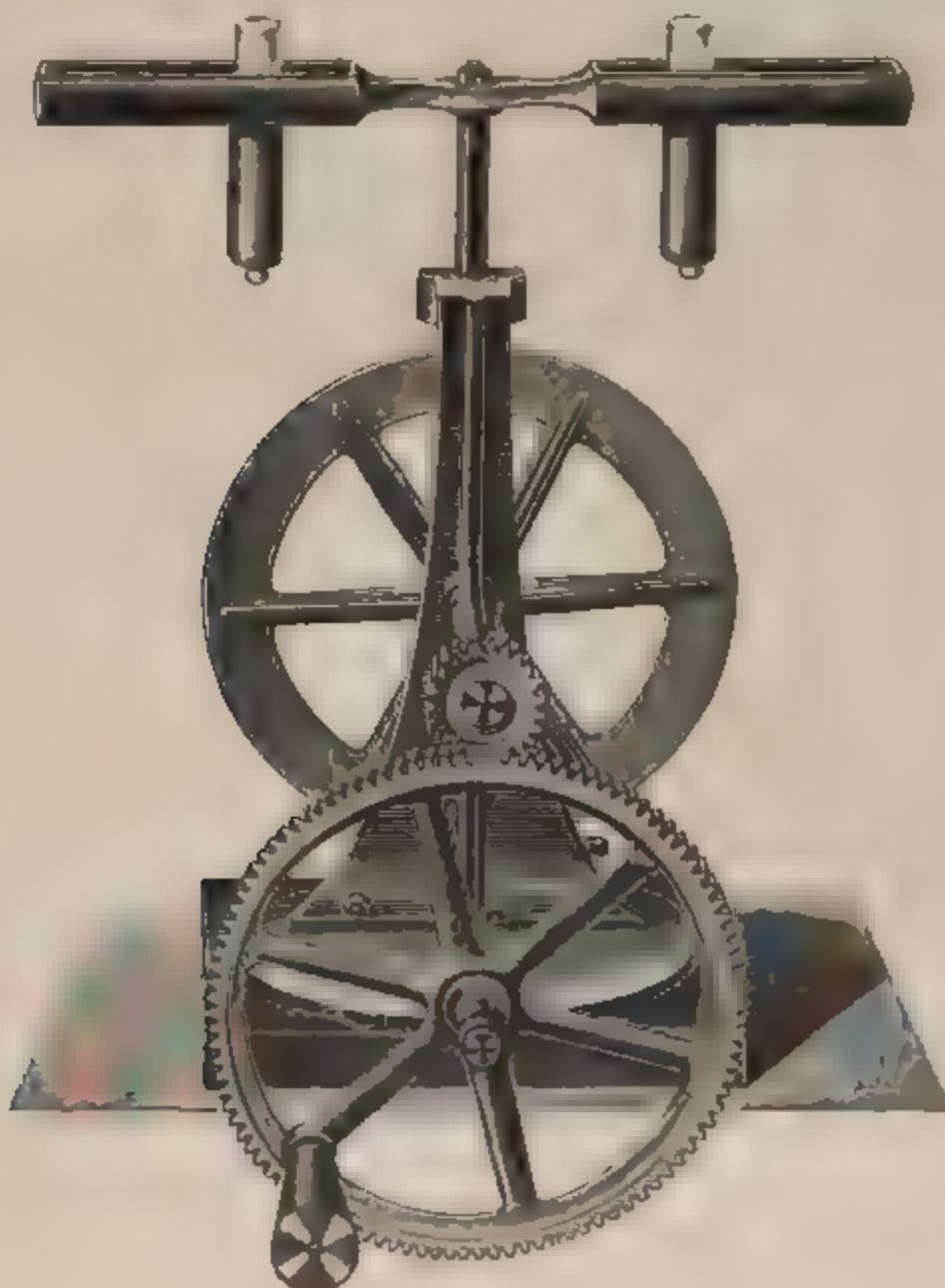


FIG. 98.—Stenbeck's Sedimentator

1. Cellular Constituents (Organised Sediment) of the Urine.

1. **Red Blood Corpuscles.** Red blood corpuscles occur in variable quantity as a morbid constituent of the urine. They may be so few in number as not to affect the colour of the fluid, and to be discernible only with the microscope; or they may be present in such abundance as to form a layer several centimetres deep at the bottom of the vessel,

and when intimately mixed with the urine impart to it a deep red colour.

The condition in which they are found is not less liable to vary than their number. They may retain their proper form, or they may appear as pale yellowish rings (phantom corpuscles of *Traube*). (See fig. 102.) Important inferences as to their origin are to be drawn from the number and character of the red blood-cells in the urine. Such cells may be derived from the urethra, the bladder, the ureters, the renal pelvis, or the kidneys themselves.

When they are intimately mixed with the urine, in such a way that though present in large quantity and deeply tinging the fluid, they do *not* form a sediment after many hours' standing, it may be inferred that the haemorrhage took place in the substance of the kidney or in the renal pelvis or ureters. If, under these circumstances, they are seen with the microscope to be profoundly altered, having lost their colouring matter and presenting the appearance of pale yellow rings, the further conclusion results that the blood was effused in the kidney itself, and the symptom points to acute nephritis or to a fresh exacerbation in the course of chronic nephritis. When the blood-cells appear as comparatively few, attenuated, and washed-out rings, they may have originated in congestion of the kidneys or in miliary tuberculosis of those organs, and the diagnosis of these conditions may be established on this ground—in conjunction, of course, with other symptoms.

It is more difficult to decide from the character of the blood passed whether the lesion has taken place in the renal pelvis or the ureters. To arrive at a just conclusion on this point, a careful examination must be made for the other occasional organic constituents of the urine, as epithelium, casts, &c., and the diagnosis will rest in a large measure upon their character.

When blood is present in considerable quantity without being intimately blended with the fluid, it is derived in the majority of cases from the bladder. Intermittent haematuria, attended with severe pain, is caused by calculi or tumours in that organ. Haemorrhages into the kidney occur in haemophilia³⁸ and leukæmia. Transient haematuria is found to occur in many morbid states, without any evidence as to its cause being disclosed post-mortem.

2. Leucocytes.—The urine contains isolated leucocytes in health. It is only when they occur in greatly increased quantity, or in conjunction with other cellular elements of a pathological character (casts), that their presence attains any serious import. They are usually unaltered in form, but sometimes, and especially in alkaline urine, they swell up, become glossy and homogeneous, and their nuclei disappear, but can again be made visible with acetic acid. Occasionally they enclose

much fatty matter, and this chiefly when they are derived, not from the urinary passages themselves, but from the bursting into them of a slowly formed abscess of some neighbouring organ, as the rectum or prostate.

Leucocytes are occasionally seen to present protoplasmic processes. This happens when the secretion possesses a feebly alkaline reaction.

The leucocytes of the urine may be derived from the substance or the pelvis of the kidney, the ureters, the bladder, the urethra, or, as already mentioned, from the rupture of an abscess into some part of the urinary passages.

These bodies may form a compact layer of sediment several centimetres deep in purulent catarrh of the bladder, and a similar deposit of pus has been found in the urine in cases of acute infectious urethritis (gonorrhœa) (*v. Jaksch*). Such pus is thick and viscid, and the constituent leucocytes are usually much altered in character (*vide supra*).

Pus-cells occur in the urine in considerable quantity in inflammation of the ureters and in pyelitis, but in this connection they are never so abundant as in cystitis, and they are commonly deposited in the form of a flocculent sediment of a slimy translucent appearance, which, when looked at through the microscope, is seen to consist of a more or less dense aggregation of leucocytes. The distinction as to the nature of the sediment is not, however, altogether characteristic, nor is it in either case invariable. With this qualification, it may be recommended as a point worth attending to in the differential diagnosis of cystitis and inflammation of the pelvis and ureters.

In renal disease there are but few leucocytes to be found in the urinary sediment, unless in the rare cases where a renal abscess discharges directly into the large tubules or the pelvis of the kidney.

Care must be taken in the case of women to ascertain whether pus found in the urine may not have been derived from the vaginal secretion. In blennorrhœa a considerable quantity of pus may find its way from this source into the urine.

When pus in great quantity appears suddenly in the urine (pyuria), it points to the opening of an abscess into the urinary passages; but two cases are on record (*r. Jaksch*) where no such causes could be assigned, and these constitute rare but remarkable exceptions to the principle laid down. They depended, doubtless, upon unusual conditions favouring diapedesis.

In these two cases the patients—a boy of six and a girl of thirteen years—both suffered from pulmonary phthisis, and the accident referred to occurred during the last week of life. The purulent sediment of the urine contained no bacilli, and the post-mortem failed to disclose any condition to which the discharge might be referred.

The fact is of importance, since it shows that a purulent deposit may appear in the urine apart from the causes mentioned above. *Glaser*³⁹ has shown that the urine of healthy persons may contain great quantities of leucocytes as a result of alcoholic excess.

Leucocytes can generally be recognised by simple inspection with the microscope; but should any doubt arise as to their nature in a particular case, it may be at once resolved by the addition of a little iodo-potassic-iodide solution. With this reagent the leucocytes stain a deep mahogany-brown (glycogenic reaction), whilst the forms of epithelium, with which they are occasionally blended, and which may be confounded with them, assume a light yellow colour.

*A. Vitali*⁴⁰ recommends the following test:—The suspected urine, if alkaline, is acidulated with acetic acid and passed through a thick filter. The deposit on the filter is then treated with a little guaiacum tincture which has been kept in the dark. If pus be present, the inner surface of the filter takes a deep blue tint. *E. Frank*, who has employed this test in the author's clinic, reports in very high terms of its efficiency. The result is obtained even with a small proportion of leucocytes in the urine.

3. Epithelium.—The slight cloud which ordinarily forms in healthy urine contains a number of epithelial cells. These are for the most part of the squamous variety, but amongst them are also some smaller forms, which are derived chiefly from the mucous surface of the renal pelvis and ureters, and very rarely from the substance of the kidneys.

In addition to these, there are to be seen in every specimen of urine a considerable number of uninuclear polygonal cells, and similar round cells, which are their earlier form. These belong to the meatus and prepuce, and in women to the vagina. Their presence in comparatively small numbers has no special significance; but when they occur in excess, they indicate a catarrh or catarrhal irritation of the parts from which they are derived. A form of epithelium, consisting of oblong cylindrical cells, diminishing in size towards their attached extremity, and with well-defined borders, comes from the surface of the male urethra (*Bizzozero*).

It is very difficult to distinguish between the epithelial cells which are derived respectively from the bladder, ureters, and renal pelvis. *Bizzozero*⁴¹ maintains that the cellular type is the same for all these parts, and *Eichhorst*⁴² agrees with him.

It follows that a particular affection in one of these situations can hardly be localised by the character of the urinary epithelium. The cells in each case have this in common, that they are smaller than those already described. Those that come from the superficial layers of the mucous membrane are polygonal or elliptical in shape. They

usually have a single large nucleus, and their protoplasm is apt to be very granular. The cells which are derived from the middle and deeper layers are more oval in shape, often irregular and conical, and furnished with one or two long protoplasmic processes (fig. 99, *b*, *b'*, *b''*). They also have a single nucleus of large size, and their substance is clearly granular. Like *Bizzozero* and *Eichhorst*, *v. Jaksch* has been unable to find any morphological distinction among these cells, by which their origin in the bladder, ureter, or pelvis can be known; nevertheless, he is of opinion that certain inferences may be drawn from their number. Given a disease of one of these parts, it may be assumed to involve the ureter alone when the epithelial forms which we have been considering are very few. When in greater quantity and superimposed upon one another like the tiles of a roof, they probably come from the pelvis; and when in very great abundance, they indicate cystitis. These points should not be strongly insisted upon, but, taken in conjunction with other symptoms, they should carry some weight in a differential diagnosis.

The general pathological condition to which the presence of these cells in excess is, in any case, to be ascribed, is that of irritation or inflammation of the mucous surface of the bladder, ureter, or pelvis of the kidney. If the reader will refer to what has been said of the leucocytes in the urine in connection with these diseases, he will see that the character of the sediment in the two particulars taken together, and read in conjunction with the clinical symptoms, will suffice very accurately to discriminate cystitis from inflammation of the pelvis or ureter.

Much importance attaches to the appearance in the urinary sediment of epithelial cells derived from the mucous lining of the tubules of the kidney. Under normal conditions they are distinguished by their smaller size from the forms just considered, or at any rate from such as belong to the middle and deeper layers of the urinary mucous membrane. They are polyhedral in shape, and finely granular, with comparatively large oval nuclei and nucleoli. They occur separately or cohering in groups (fig. 99, *c*, *c'*, *c''*, *c'''*), and in the latter case may display the cylindrical arrangement (epithelial casts, fig. 101). They are often to be seen, singly or several together, on the surface of the casts to be described later (fig. 110, *c*).

The cells of kidney epithelium exhibit remarkable deviations from the normal type. They are sometimes hard, tough, and glossy, like the obsolete cells of the intestine described by *Nothnagel* (p. 199). They occasionally contain fatty globules in greater or less profusion (fig. 99, *d*, *d'*), and again cells are seen overlying the surface of casts (fig. 109, *a*), which may conform in shape to the type described above, but consist entirely of fatty matter (see also fig. 109, *c*).

In the convalescent stage of acute nephritis (of scarlatina and erysipelas), small round cells with an eccentric resting nucleus are frequently to be seen, and these are doubtless the young kidney-cells formed in the process of repair within the tubules (*v. Jaksch*).

The detection and discrimination of the various forms of kidney epithelium in the urine are matters of the highest importance in diagnosis. The presence of epithelium is always a sign of renal disease, and usually of inflammation. In addition to this, and where the coincident symptoms indicate a nephritis, the character of the epithelium will enable us to form a probable opinion as to whether the inflammatory process is accompanied with degeneration of the kidney substance. Thus where the cells are found loaded with fat, the autopsy will most likely disclose a fatty condition of the renal tissues; and the detection

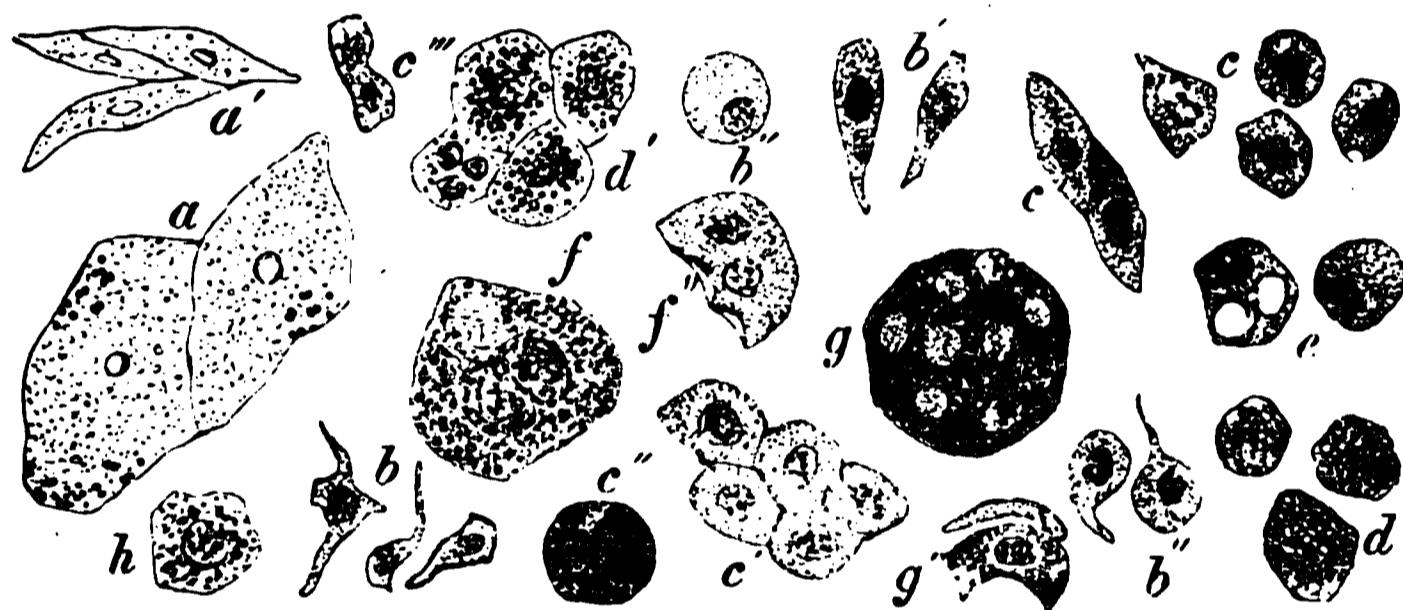


FIG. 99.—Urinary Epithelium (eye-piece III., objective 8A, *Reichert*), collected from thirty different specimens.

a, a'. Squamous epithelium from the urinary sediment.
b, b', b''. Epithelium from the bladder.

c, c', c'', c'''. Epithelium from the kidneys.
d, d'. Fatty epithelium from the kidneys.

e-h. Epithelium from the bladder.

of the obsolete cells described above points with some degree of certainty to an amyloid degeneration of the organ.

It is needless to add that a diagnosis can never be based upon such appearances alone, but must in all cases be controlled by the clinical symptomis and the other manifestations of the urine.

4. Casts.—The subject of urinary casts, which will now engage our attention, is one of very great interest. Casts were first seen in the urine by *Vigla*, *Quevenne*, and *Rayer*,⁴³ in France, and almost simultaneously in Germany by *Simon* and *Nasse*.⁴⁴ *Henle*⁴⁵ discovered them in the urine of a dropsical patient, and afterwards in the renal tubules both of health and disease. *Glaser*,⁴⁶ working in the author's clinic, has ascertained that the recent urine of healthy persons, while free from albumin, often contains casts, and that a slight toxic influence (alcohol) is often

sufficient to determine their presence. *Rorida*⁴⁷ has contributed the most ample information concerning urinary casts. These bodies are subject to very great variety as to their number when present, their form, and the import which attends their manifestation. It must be premised that they have been found in urine which is entirely free from albumin, and even from every other morbid product. Thus *Nothnagel*⁴⁸ has seen them in the urine of patients with jaundice, which at the same time contained no albumin; *Burkart* and *Fischl*⁴⁹ likewise observed them, in the absence of albumin, in cases of severe inflammatory affections of the stomach and intestine. *Radomyski*⁵⁰ has observed casts in albumin-free urine, in connection with disturbances of the circulation. *Kohler*⁵¹ found cylinders, cylindroids, and renal epithelium in the urine, in cases of intestinal disturbances of an acute or chronic nature, as well as in cases of obstipation, without these symptoms being always accompanied by albuminuria. Hence it follows that the presence of casts in the urine does not by itself imply disease.

Urinary casts may be conveniently divided into two chief classes, viz., *Unorganised* and *Organised*.

(a.) *Unorganised Casts*.—These are formed of crystals. They are pathologically of little consequence. Those that have been described consisted of urates (fig. 100) and hæmatoidin; and as yet they have been found only in infants, and in cases of gout and renal congestion. If healthy urine be concentrated in vacuum at a low temperature, 37°–39° C., casts may be observed which consist of acid urate of soda (*Leube*).⁵²

Perhaps we ought to include under this heading some of those bodies which are at present classed together under the general name of "detritus" casts.

(b.) *Organised Casts* consist of cellular elements, or the products of their transformation.

They may be subdivided into three groups: (1.) Those which consist of cells (red blood-corpuscles, leucocytes, and epithelial cells). (2.) Those which consist of the products of cellular change. (3.) Those so-called hyaline casts, whose origin is still a subject of dispute, but which are sufficiently distinguished from the others, both clinically and morphologically.

(1.) The first group includes:—

- (a.) Casts formed of red blood-corpuscles (fig. 102).
- (b.) Casts formed of leucocytes (fig. 103).
- (c.) Epithelial casts (figs. 101 a, b, and 104 a, b).
- (d.) Casts consisting of colonies of bacteria.

(2.) Under the second group we distinguish:—

- (a.) Granular, (b.) waxy, and (c) fatty casts.

(3.) The group of hyaline casts may be subdivided according as its members are simply hyaline, or, in addition, coated with certain substances, amongst which kidney epithelium, red and white blood-corpuscles, bacteria, and various forms of crystals may be enumerated.

The cylindroids of *Thomas* should perhaps be included in this group.

The number of casts occurring in a specimen of urine, and the length and breadth of each, is subject to very great variety. The latter fact is sufficiently seen in the accompanying figures.

(1.) Perfectly formed cellular casts appear in the urine only under such circumstances as cause the renal tubules to become crowded with red or white blood-corpuscles, or bring about the separation of the renal epithelium in the entire circumference of a tubule. When this occurs, the casts formed are forced onward by the flow of the fluid secreted behind them, and are ultimately discharged from the bladder.

Fig. 104 (*a* and *b*) shows a rare specimen of casts, consisting of renal



FIG. 100.—Cast of Urates, from a case of Emphysema (eye-piece III., objective 8A, Reichert).

epithelium and leucocytes from the urine in a case of nephritis with oliguria and uræmia.

The clinical significance of these casts is very great. *They always imply an affection of the kidney, and their presence alone suffices to establish the existence of acute nephritis, or the supervention of a fresh paroxysm in that disease.* When found in small numbers the kidney is probably but slightly diseased. When, on the other hand, the urine holds abundance of these casts, the fact is ample evidence of inflammatory changes in the organ. Figs. 101, 102, and 103 represent specimens of cellular casts formed in different proportions of the several cellular elements which enter into the formation of these bodies.

Casts are sometimes found which consist almost entirely of masses of micrococci, but these have quite a different significance. Fig. 113, *d*, represents such a one. Morphologically, they bear a close resemblance to the granular casts presently to be described, but are distinguished

from them by their resistance to powerful reagents, such as caustic potash and nitric acid. They may also be known by their opacity and



FIG. 102. Epithelium cast, from a case of chronic nephritis. *a*, fully differentiated; *b*, in part granular (eye-piece III, objective 8A, Reichen).

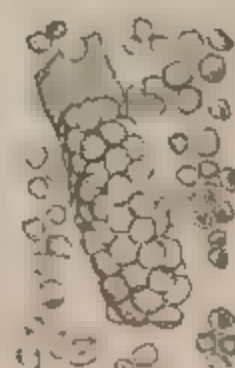


FIG. 102.—Cast of shrunken blood-corpuscles in part metamorphosed into neutrophil leucocytes, from a case of acute nephritis (eye-piece III, objective 8A, Reichen).



FIG. 103.—Cast formed of leucocytes, from a case of acute nephritis (eye-piece III, objective 8A, Reichen).



FIG. 104.—*a* and *b*. Rare forms of casts composed of leucocytes and epithelium, from a case of chronic nephritis which began with oliguria and uremic paroxysms (eye-piece III, objective 8A, Reichen).

the grey colour which they display, as well as by the remarkable uniformity of their substance (*Martini*).¹³

Casts of micrococci in the urine are a matter of very grave significance. They imply, as a rule, the existence of septic embolism of the

kidney. They may also arise from the extension upwards of a septic pyelitis (pyelo-nephritis).

The author⁵⁴ has seen a number of casts formed of minute bacilli in

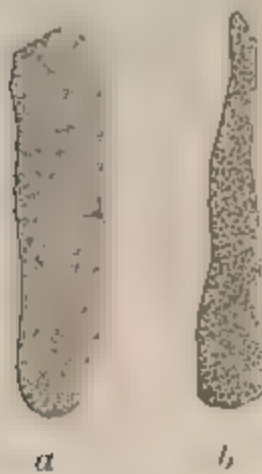


FIG. 105.—*a* and *b*. Granular cast in chronic nephritis (eye-piece III, objective 8A, Reichert).



FIG. 106.—*a* and *b*. Granular cast in acute nephritis (eye-piece III, objective 8A, Reichert).



FIG. 107.—*a* and *b*. Granular cast in chronic nephritis (eye-piece III, objective 8A, Reichert).

the urine of a boy after a few days' illness with acute nephritis. Their formation was transitory.⁵⁵

The specimen represented in fig. 113, *d*, was from fermenting diabetic urine, and had no connection with the symptoms.

(2.) The members of the second group, as we have seen, are granular, waxy, and fatty casts.

(a.) **Granular Casts** vary much in dimensions. They are most frequently seen in a fragmentary state, but are occasionally of perfect form. Their borders, which are usually well defined, are often sinuous in casts of some length (fig. 107, *a* and *b*). In the latter case, too, they are somewhat concave at the extremities; but when very short and

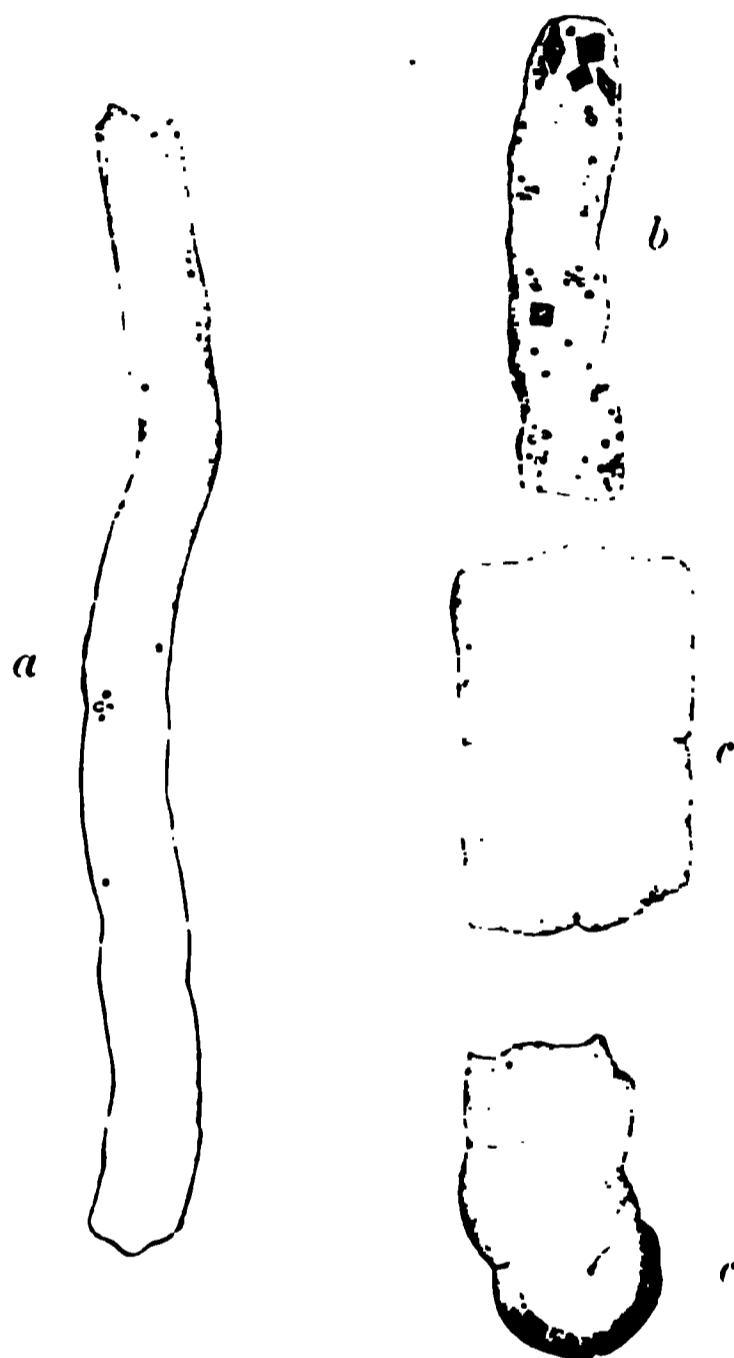


FIG. 108.—Different Forms of Waxy Casts: *a*. with a coating of urates; *b*. waxy casts covered with crystals of oxalate of lime; *c*. fragments of waxy casts (eye-piece III., objective 8A, *Reichert*).

broken, they present a zigzag ending. Their constitution is very variable. Sometimes they consist of fine particles, which can be distinguished only by a high power of the microscope (fig. 105, *a*), as Zeiss's objective F. In other cases they are coarsely granular, and the constituent particles can be readily made out with Hartnack IV. (fig. 106, *b*). In colour, too, they manifest considerable differences. They may be of all shades, from pale yellow to a reddish brown. They are occasionally coated with leucocytes, fatty globules, and needles of fatty

crystals (figs. 107, *b*, and 109, *a*, *b*). These distinctions of character are not known to correspond to special derangements of the kidney.

Granular casts sometimes exhibit a transition form of epithelium (fig. 101, *b*), and it is probable that they usually originate in the degeneration of the blood and epithelial casts already described. This theory was first stated by *Rindfleisch* and *Langhans*.⁵⁶

The presence in considerable quantity of granular casts in the urine indicates an inflammatory condition of the kidneys. They have been found (*v. Jaksch*) as an exceptional constituent of the secretion in cyanotic induration of the kidney, and especially when the latter condition was associated with nephritis (secondary nephritis). These bodies may be regarded almost equally with cellular (epithelial) casts as a certain indication of nephritis.

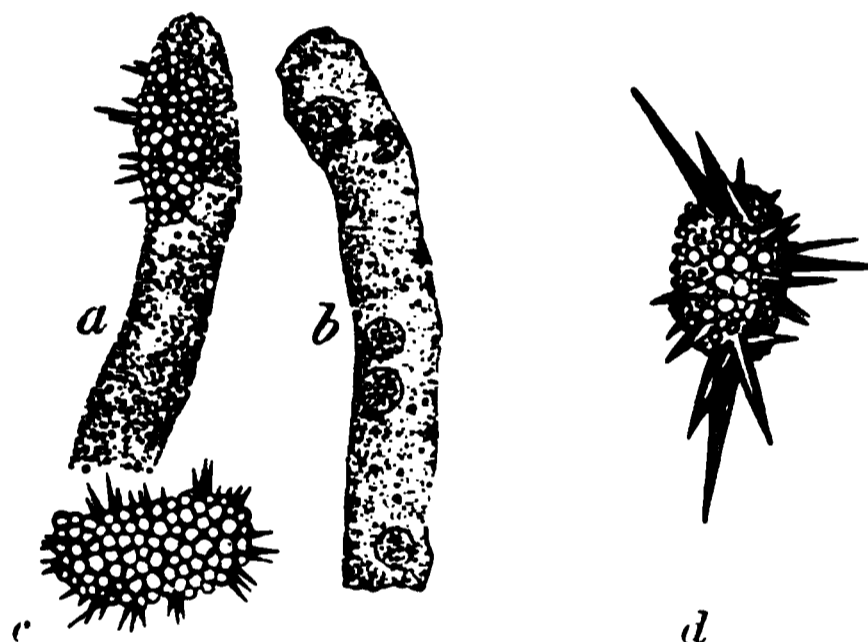


FIG. 109.—*a*. Granular cast beset with fatty globules and fat crystals; *b*. granular cast covered with leucocytes; *c*. and *d*. fatty casts; from a case of nephritis and large white kidney (eyepiece III., objective F, Zeiss).

(b.) **Waxy Casts.**—These casts attain a greater length than the others, and when comparatively perfect they may be seen to be segmented like a tapeworm. They also occur in shorter fragments of relatively great breadth. Under the microscope they are homogeneous and refractive, and often bear upon their surface fatty globules, separately or in confluent masses, white and red blood-corpuscles, fungi, and crystals of various kinds.

Their constitution has not been definitely ascertained. It would appear to be very complex, and that such casts result alike from the breaking down of epithelium and from the exudation of occasional products (fibrin, amyloid material) into the renal tubules. Their number also is very inconstant.⁵⁷

They are not characteristic of any special disease. They are found in acute and chronic nephritis, in contracted granular and in amyloid

kidneys. They frequently exhibit the amyloid reaction with methyl-violet and iodo-potassic-iodide solution; and this in the absence of amyloid degeneration of the kidney. Moreover, the reaction is not obtained in some cases where this condition exists, and consequently no inference can be drawn from its manifestation.

(c.) **Fatty Casts.**—Fatty globules are found upon the surface of granular casts (fig. 109, *a*); but they also form by themselves short, powerfully refracting cylinders, which often are beset all over with needles of fatty crystals (fig. 109, *c, d*).

These casts and fatty crystals were first pointed out by *Knoll*. They are most commonly associated with subacute and chronic inflammations of the kidney of protracted course, with a tendency to fatty degeneration of the renal tissues (*v. Jaksch*). Consequently their detection affects the prognosis unfavourably.

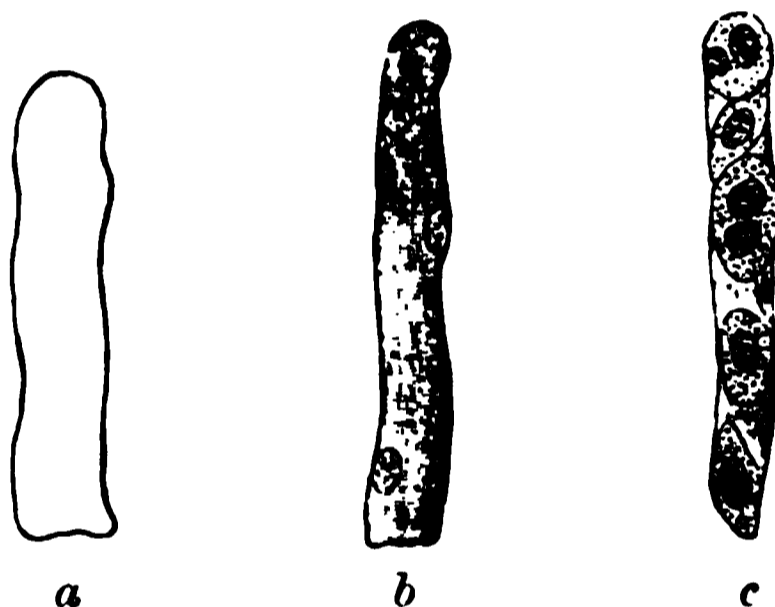


FIG. 110.—Hyaline casts; *a*. hyaline cast; *b*. hyaline cast coated with leucocytes; *c*. hyaline cast covered with kidney epithelium; from a case of chronic hypertrophic hepatitis with jaundice (eye-piece III., objective 8A, *Reichert*).

Post-mortem examination has shown that they form most frequently in cases of large white kidney. In some instances where they were present, however, the organ was found to be more or less contracted; but when this was so, it was invariably far advanced in fatty degeneration. The crystals which beset fatty casts are not formed exclusively of fat, but probably to some extent of lime and magnesia salts of the higher fatty acids and allied compounds, for they are *not* all soluble in æther. They have their origin doubtless in fatty degeneration of the renal epithelium.

3. Hyaline Casts.—Hyaline casts are for the most part pale and delicate bodies, of varying length and thickness, and more or less abundant in different specimens. They cannot, as a rule, be detected without being previously stained. Their pathological significance differs greatly according as they are coated with certain substances or not.

In some diseases, whilst the urine contains no albumin, very pale hyaline casts are found in small numbers in the sediment. *Nothnagel*⁵⁸ has met with such in albumin-free urine in cases of jaundice, and *Henle*⁵⁹ in others where the kidneys were quite healthy. Their presence, therefore, has no necessary connection with renal disease. The author has repeatedly been able to recognise hyaline casts in the urine in affections whose subsequent course altogether excluded the possibility of diseased kidney; and *Huppert*⁶⁰ has shown that the urine voided after an epileptic paroxysm may contain both albumin and hyaline casts in the absence of any kind of inflammation in that organ. According to *Leube*,⁶¹ hyaline casts are seldom met with in urine free from albumin.⁶² Nothing more need be said to enforce the precept that the existence of renal disease, or at all events of nephritis, must not be hastily concluded from the mere presence of hyaline casts.

But when these casts bear a coating of other substances upon their surface the case is altogether different. In nephritis we sometimes find in conjunction with other casts cylinders of hyaline substance which are covered with epithelium (fig. 110, *c*), unaltered or loaded with fat, leucocytes (fig. 110, *b*), and red blood-corpuscles.

In severe cases of jaundice depending upon disease of the liver, as in secondary carcinoma of that organ complicated with nephritis, colourless hyaline casts are formed in the urine, and overlying their substance are golden-yellow cells of kidney epithelium, which colour red, changing to blue in presence of nitric acid.

Urates are seen deposited upon such casts in cases of congested kidney; and other bodies, as oxalate of lime and bacteria, may similarly occupy their surface.

This is perhaps the place where mention should be made of the so-called cylindroids (fig. 111). These are long, ribbon-like bodies, resembling casts, which were first discovered by *Thomas*⁶³ in the urine of scarlet fever. They occur also in nephritis, cystitis, and renal congestion, and *Bizzozero* has found them in healthy urine.⁶⁴ They are not, therefore, characteristic of kidney disease. They are observed most commonly in the urine of children, which may or may not also exhibit albumin, in the absence of other symptoms. *Pollak* and *Török*⁶⁵ have noted these occurring together with abnormal excretion of urates.

With regard to the mode of origin of hyaline casts and cylindroids, *Rovida*⁶⁶ suggests that they are the products of secretion by the epithelium lining the urinary tubules, and his view is borne out by the experiments of *Pollak* and *Török*. In this way it is possible to account for their manifestation in the absence of severe renal affections. At the same time it must be borne in mind that the experiments which *Ribbert*⁶⁷ made some years ago upon animals point to the conclusion that hyaline

casts may result directly from the exudation of albumin within the tubules.

Method of Examination for Casts.—The urine should be treated, when necessary, with antiseptic agents (*v. supra*). The author has found, by several years' experience, that the addition of a little powdered camphor affords one of the best means for preserving urine from decomposition. Such an addition leaves the cellular elements entirely unaltered. It should be covered and allowed to stand for several hours (or the proceeding may be expedited by the use of the sedimentator), and some of the sediment which has then fallen should be removed with a pipette for examination on a slide.

The casts belonging to the first two classes can usually be recognised without special preparation. Simple hyaline casts may need to be stained to render them visible, and this may be done best in their case by the use of a drop of dilute solution of iodine and iodide of potassium. Other staining fluids may be used for colouring the different varieties of casts. These are picro-carmine, gentian-violet, eosin, acid haematoxylin solution, safranin, Bismarck-brown and methylene-blue. It should be mentioned, however, that the staining properties of casts vary greatly, and that some which are morphologically similar will behave very differently in solutions of the substances enumerated here. *Rieder*⁶⁸ recommends the employment of the dye-stuff Soudan III, by which the fatty globules in particular are rendered distinctly visible. Personal experience confirms this report.

For the investigation of these staining properties the sediment should be washed with the normal saline (75 per cent. NaCl) solution (*Knoll*⁶⁹), and care should be taken that the dyes employed are sufficiently diluted.

[For the preservation of urinary casts *Harris*⁷⁰ advocates the use of a solution made as follows: potassium acetate 60 grms., chloro-



FIG. 111.—*a* and *b*, Cylindrallia from the urine in congested kidney (eye piece II objective 8A, *Steckert*).

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FIG. 111.—*a.* and *b.* Cylindroids from the urine in congested kidney (eyepiece II., objective 8*A*, *Reichert*).

form 10 cc., distilled water 1000 cc. The fluid is placed in tubes specially made. These are closed by an indiarubber stopper at the broad upper extremity, and open below at a fine drawn point. The urinary sediment, obtained under careful precautions of contamination, is dropped in from above, allowed to settle, and finally expressed at the open point, received in the depression of a glass slide, covered with a slip, and cemented. Such preparations are said to be permanent.]

Chemical Constitution of Renal Casts.—*Rovida* is still the classical authority upon this subject. According to him, the characteristic property of hyaline casts is their solubility in weak mineral acids. This property they possess in common with cylindroids.

Waxy casts in their behaviour with chemical agents resemble albuminates, but they also exhibit reactions which sufficiently distinguish them from the latter substances. It would appear in general that the substance of which casts are formed is not a proteid, but some derivative of one. In this conclusion *Rovida* had been long anticipated by *L. Mayer*.⁷¹ *Knoll* has also satisfied himself that the substance of which renal casts are composed is not identical with any of the proteids with which we are familiar, as acid albumin, albumin, albuminate, albumose, globulin, fibrin, mucin, or peptone.

5. Spermatozoa.—These are pear-shaped bodies, about 50 μ in length. Of this, the head occupies 4 to 5 μ , and the remainder is a long tail which tapers towards the extremity (see fig. 152).

Spermatozoa are found in the urine of men after coitus, involuntary emissions, e.g., of epileptic paroxysms (*Huppert*⁷²), and masturbation. They may also occur in the urine of women when passed directly after connection (see Chapter IX.).

6. Fragments of Tumours.—It rarely happens that fragments of a tumour are found in the urine. The author has never yet by their aid been able to diagnose a growth in the kidney. On the other hand, a carcinoma of the bladder, or a tumour of some other organ, as the vagina or rectum, which has burst into the bladder, may betray its character by imparting its constituents to the urine. Thus a pigmented tumour may be known by the detection of melanotic particles in the sediment; but in other cases, as where cancer-cells are mixed up with ordinary epithelium, it is more difficult to base a diagnosis upon such appearances, and this can only be done in conjunction with the other symptoms.

Tumours of considerable size (polypi, &c.) have been known to pass with the urine. An observation by *Heitzmann*⁷³ shows that it is sometimes possible to diagnose renal tumours by a microscopic examination of the urine.

7. Parasites.

1. Fungi.—Adopting the same classification as before, we shall divide the fungi of the urine into moulds, yeasts, and fission-fungi, and we shall also consider them under the two headings of pathogenic and non-pathogenic organisms.

(a.) *Non-Pathogenic Fungi*.—Fresh healthy urine is free from fungi (*Leube*⁷⁴) ; but when allowed to stand for some time, it becomes crowded with these organisms.

All three classes of fungi may be represented in the urine. It must be mentioned, however, that where ammoniacal fermentation is in progress, as a rule only fission-fungi, and perhaps a few yeast-cells, are to be found. Moulds are, under normal circumstances, a very rare manifestation in decomposing urine ; but in that of diabetes, when the alcoholic fermentation of sugar has ceased, they make their appearance in great quantities, floating in a layer of upwards of a millimetre in thickness on the surface of the fluid, to which they impart a disagreeable mouldy smell. The urine is at the same time turbid with yeast-



FIG. 112.—*Micrococcus ureæ* from the surface of a Normal Urine undergoing Ammoniacal Fermentation (eye-piece III., objective 8A, *Reichert*).

fungi and bacteria, and its appearance alone may be conclusive as to the abundant excretion of sugar.

The development of yeast-fungi in large numbers is a sure sign of sugar in the urine, and their detection will serve to suggest this condition where it has been previously overlooked.

The microscopical character of fermenting healthy urine is subject to great variety. The transformation of urea into carbonate of ammonia is most likely effected through the agency of several forms of fungi (*Miquel*, *v. Jaksch*, *Leube*, *Billet*, *C. Flügge*, *v. Limbeck*⁷⁵), but most prominent are the micrococci. Of these, the *Micrococcus ureæ* (fig. 112) may be seen in almost pure cultivations upon the surface of the fluid. This micro-organism forms in long chain-like series, and is of comparatively large size. In addition to these, there are rod-shaped bacteria of all sizes and forms, and, as occasional manifestations, certain long spiral bacilli with large spores, and cocci, which group themselves into globular masses of dark colour and varying size (fig. 113, g). *Sarcina* is also found in the urine. It is smaller than that which forms

in the stomach, being in point of size comparable to the sarcina of the lung (see p. 125). Moreover, according to *Fr. Hofmeister*⁷⁶ the normal urine of healthy men always contains germs.

(b.) *Pathogenic Fungi*.—When recently voided urine is found to contain a profusion of fungi, the condition is in every case serious, because these fungi—even though not specifically pathogenic—may give rise to serious trouble by promoting decomposition within the bladder. The circumstances which influence the development of non-pathogenic fungi in fresh urine have been made the subject of repeated investigations (*Roberts, Schottelius, Reinhold, Barlow*⁷⁷). The matter is still surrounded with a great deal of uncertainty, and it is convenient provisionally to distinguish the condition as idiopathic bacteriuria. According to the researches of *Schottelius* and *Ross*,⁷⁸ it is unattended with any morbid symptoms. In the case of a patient who had suffered for a year from gonorrhœa and cystitis, and who said that he had never had a catheter passed, the author found the urine turbid and ammoniacal, and pro-

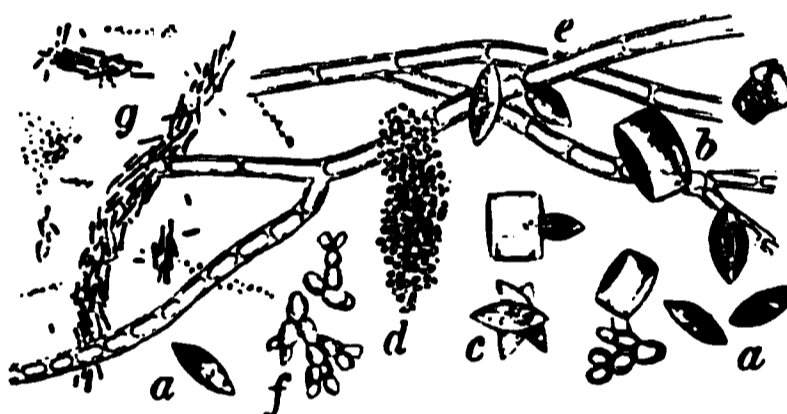


FIG. 113.—Sediment from Fermenting Diabetic Urine with Casts of Micrococci.

a, b, c. various forms of urates; d. micrococci in form of casts; e. moulds; f. yeast-fungi; g. bacilli and micrococci (eye-piece III., objective 8A, *Reichert*).

liferating crowds of micrococci of every description. This continued long after the cessation of symptoms, and was attended with pain.

The diagnosis of idiopathic bacteriuria must be made with great caution. The author had the opportunity of observing another case, apparently analogous to the one described above. From a communication recently made by the patient, however, the symptom is sufficiently explained by the appearance in the interim of a prostatic abscess.

Bacteriuria is very often observed after the use of unclean (non-sterilised) catheters.⁷⁹ Often (although not always) cystitis supervenes as a result of this. Very interesting communications on the subject of the decomposition of urine by bacteria have been made by *Crämer* and *Albertoni*.⁸⁰

Pathogenic fungi occur in the urine in connection with certain specific diseases, as erysipelas, relapsing fever, septic processes, typhoid, and tuberculosis, and it is a matter of great importance to be able to detect them.

In introducing this subject a few general remarks are called for:—

In the first place, there can be no doubt that large numbers of sufficiently characteristic micro-organisms may be present in the freshly voided urine of infectious diseases, especially when it also contains albumin and casts (*Kannenberg, Litten, R. v. Jaksch*).⁸¹ In one such disease, namely, erysipelas, the author's experience has been that, in all cases where the typical symptoms of acute nephritis supervened, the urine contained a profusion of fungi which were indistinguishable in their form from the *Streptococcus pyogenes* or *erysipelatos* (*Fehleisen*).⁸² The urine was nearly always turbid, and while still quite fresh exhibited these bodies in moniliform arrangement. In these cases it always happened that the bacteriuria and nephritis disappeared with the cessation of the erysipelas.



FIG. 114.—Tubercle-Bacilli from Urinary Sediment in a Case of Tuberculosis of Urinary Organ (eye-piece III., objective $\frac{1}{2}$, oil immersion, *Reichert*).

That it was a true nephritis which terminated favourably in all these cases was sufficiently shown by the results of microscopical and chemical examination, which disclosed the presence of albumin, blood casts (of Groups I. and II.), renal epithelium, and numerous leucocytes in the urine.

It has been already mentioned that in septicæmic processes the urine has repeatedly been seen to hold cylindrical bodies, whose chemical properties showed them to consist of micrococci (*Martini, Litten, Senetz*).⁸³ *Weichselbaum*⁸⁴ has found specific micrococci in the urine in ulcerative endocarditis; *Lustgarten* and *Mannaberg*⁸⁵ cocci in acute nephritis; and *Letzerich*⁸⁶ bacilli in the "primary" nephritis of children. *Mircoli*⁸⁷ also determined the existence of pneumonococci-like forms in the urine of children suffering from this disease. Further, *Neumann*⁸⁸ has found the typhoid-bacillus in the urine in six out of twenty-three cases investigated, *Wright* and *Semple*⁸⁹ in six out of seven, and *Karlinski* and *Konjajeff*⁹⁰ have obtained cultivations of the bacillus from typhoid urine in the earliest period of the disease. *Philipowicz*⁹¹ has recognised the tubercle-bacillus and the bacillus of glanders in urine.

The spirilla of relapsing fever (see Chapter I.) occur very rarely, and only when haemorrhage takes place in the kidney during the period of exacerbation ; but *Kannenberg*⁹² asserts that various forms of microbes are detached from the kidney in very great numbers in the exacerbations of this disease.

The recognition of tubercle-bacillus in the urine has of recent years been invested with great pathological interest (*Leube, Rosenstein, Babes, Shingleton Smith, Irsai, Benda, Kreske*).⁹³ The method of its detection is the same as that already described in connection with the sputum (see Chapter IV.). Its presence in general points to tubercular ulceration in some part of the urinary tract, and most unequivocally when the bacilli are found to be arranged in S-shaped aggregations (fig. 114), or in colonies of unmixed constitution (pure cultivations). It should be stated, however, that *Philipowicz*⁹⁴ has discovered isolated specimens of the bacillus in the urine in miliary tuberculosis, where there were no tubercular ulcers in the genito-urinary passages. The localisation of ulcerated patches must depend upon the other microscopical constituents of the urine. When these point to an affection of the kidney, we are warranted in inferring tuberculosis of that organ.

A form of caseation is known to occur in the kidney which, in its gross appearances, closely resembles chronic tuberculosis. And in this condition a careful examination, whether of the urine or of the caseous masses removed from the organ after death, will fail to reveal the specific bacillus. It would appear, therefore, that chronic non-specific inflammatory changes, such as we are familiar with in the lungs, may take place also in the kidney, and lead to destruction of its tissues.

When, in the course of pulmonary tuberculosis, the urine is found to contain albumin or pus, it will suggest to the physician the possibility of the renal complications which are known to attend the diathesis ; he will diligently examine the urine for the tubercle-bacillus, when the symptoms, after microscopical, chemical, and clinical investigation, do not find their explanation in the assumption of amyloid degeneration of the kidneys, of chronic nephritis, or cystitis.

Actinomyces may also appear in the urine in cases where the genito-urinary tract is infested with it, or when discharged thereby from other parts.⁹⁵

In searching for pathogenic fungi, it is essential that the parts about the meatus be carefully cleansed, and the urine passed into a thoroughly disinfected vessel.⁹⁶ It should then be allowed to settle, or the sedimentator may be used, and cover-glass preparations made from the sediment in the usual manner. In some cases it will be necessary to resort to Koch's method of plate-cultivation to obtain the various micro-organisms in an unmixed condition. Finally, the inoculation

of animals will resolve doubt as to the character of the specific organisms obtained.

2. Infusoria.—The author has frequently observed infusoria in the urine, but never when it was fresh. They made their appearance in all cases only when it had been allowed to stand for some time, and the fluid containing them was generally feebly alkaline. Amongst these organisms were bodies which were similar to the cercomonad already described in the chapter on *Faeces*. *Hassall*⁹⁷ has given to one of the infusoria of the urine the name of *Bodo urinarius*.

The presence of infusoria has no pathological significance. *F. Marchant*⁹⁸ found *Trichomonas vaginalis* in a man apparently after the bursting of a pelvic abscess into the bladder. *Muir*⁹⁹ has made a similar observation, and *A. Dock*¹⁰⁰ has confirmed the above by his own experience. *Budz*¹⁰¹ observed a great quantity of amœbæ in the turbid urine of a girl twenty-three years of age who was affected with phthisis

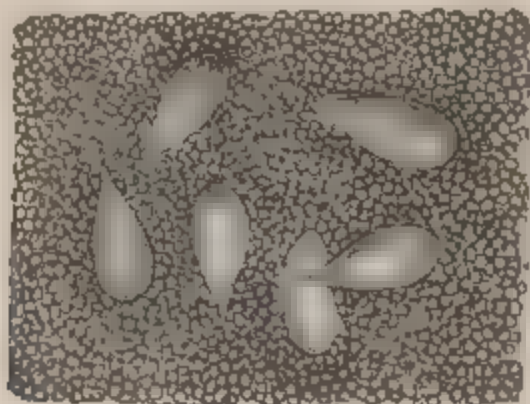


FIG. 115.—Eggs of *Distoma Haematobum* in Sediment (eyepiece II., objective C., Zeiss).

These were of larger size than the forms already described as occurring in the stools.

3. Vermes.

1. *Distoma Haematobum*.—The eggs of this parasite are often found both in the urinary passages and in the urine of inhabitants of the tropics (see Chap. I.). In such cases the latter furnishes other evidence of the parasite, as blood, and frequently abundance of fat. The condition is attended with severe burning pains in micturition.¹⁰² These are momentary, and are caused by the passage of the eggs (see Chap. I.) along the urethra, which they irritate with their sharp angles. With the last drops of urine a blood-clot is often passed. The fluid is usually clear, and contains blood-cells and pus, amongst which the eggs represented in fig. 115 are to be sought.

2. *Filaria Sanguinis Hominis*.—Lewis has detected filaria in the urine in some cases in which the blood was much infested by it (fig. 36),

and its appearance was generally accompanied with blood and pus. It is most likely this worm which causes the tropical haematuria of which *Wucherer* first sent an account from Brazil.

3. Echinococci.—The hooklets and fragments of the cysts of echinococcus occur as a very rare manifestation in the urine (*Mosler*).¹⁰³ In such cases, the cysts may have formed originally in the urinary passages—and this is very exceptional—or they may have found their way into them by rupture from a neighbouring organ. When the hooklets and characteristic membrane (fig. 63) are present, the sediment is likely also to contain red blood-corpuscles, numerous leucocytes, and a quantity of cellular débris from that part of the urinary apparatus which has been injured by the separation of the cyst.

4. Eustrongylus Gigas.—This parasite has been said to exist in the urine, but the researches of *Leuckart*¹⁰⁴ have thrown doubt upon the matter. At all events, it must be very rare. It was found by *Moccato*¹⁰⁵ in the urine of a woman—its discharge being attended in this instance with chyluria.

5. Ascarides.—In exceptional instances ascarides make their way from the intestine into the urinary passages. Their presence in the urine is usually brought about by an abnormal communication between the latter and the alimentary canal. *Scheiber*¹⁰⁶ has recently found in the urine of a woman worms which he thought had been derived from the genital organs, and he has named them *Rhabditis genitalis*. Similar observations have been reported by *E. Peiper, Westphal, and Baginsky*.¹⁰⁷

II. CRYSTALLINE AND AMORPHOUS DEPOSIT (UNORGANISED SEDIMENT).—It will be convenient in treating of the inorganic deposit of the urine to consider, in conjunction with the microscopical appearance of its constituents, some of the more characteristic of their chemical and histochemical properties.

The colour of the sediment and the reaction of the urine will often afford some indication of the nature of the deposit. Thus, if sediment of a deep red colour forms when the urine has stood for a short time, it consists mainly of urates. The colour is then due to the presence of urinary pigments, carried down by the deposit, uric acid and its salts being themselves colourless. If the deposit redissolves on the application of heat *without the addition of acid*, it is a further proof that it is composed of urates.

If, on the other hand, the urine is alkaline, and deposits a white flocculent sediment, the latter probably either consists of pus or contains a large proportion of phosphates, carbonates, and alkaline urates. Such a sediment is *insoluble* by heating, but readily so in presence of acid (acetic acid).

A third kind of sediment, of a mixed character, may be distinguished. This consists of urates and phosphates, and it forms in urine which was acid when passed, but which has become gradually alkaline from the supervention of ammoniacal fermentation.

An abundant sediment of urates belongs to the urine of fever and renal congestion, and occurs also in healthy individuals after excessive perspiration without partaking freely of water (*vide supra*). It is thought by *Mygge*¹⁰⁸ that the presence of such sediment, when it consists of uric acid, has a certain clinical significance, inasmuch as it is generally found in cases of rheumatism or renal disease.

Phosphatic sediment is apt to form in all conditions where the urine possesses an alkaline reaction when passed. It does not, however, always indicate disease, because it may be induced by the drinking of aerated water and by other means. Amongst pathological conditions dyspepsia often occasions the deposit of phosphates, and in general it may be said that phosphatic sediment is associated with chronic, as distinguished from acute, affections.

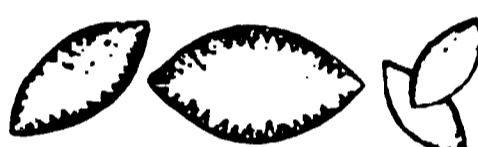


FIG. 116.—Pointed Crystals of Uric Acid from the Urine in Congestion from Emphysema (eye-piece III., objective 8 \times , *Reichert*).



FIG. 117.—Uric Acid Crystals from the Urine in Congestion of Heart-Disease (eye-piece III., objective 8 \times , *Reichert*).

The characters above referred to will serve to indicate which class of salts preponderate in the urine. A more accurate estimation of its inorganic constituents can be formed only from a microscopical and histochemical investigation of the sediment.

Such constituents may be *crystalline* or *amorphous*. Their nature will further differ according as the precipitate is an acid or an alkaline urine. We shall, therefore, consider the sediments of acid and of alkaline urine separately.

A. Sediments of Acid Urine.

(a.) Crystalline Deposits.

1. **Uric Acid.**—Uric acid occurs in crystals, which are deeply stained a brownish-yellow [or red] colour, and differ much as to their form and size. [The colour is due to uroerythrin, which has a great affinity for the crystals.] They are sometimes large and thick, in shape like a whetstone (figs. 113, a, and 116), and then commonly exhibit a dark nucleated centre; sometimes they are longitudinally striated spicules (fig. 117) and sometimes again rhombic tables (figs. 113, b, and 116) with rounded

angles. They may appear separately or in masses. Their shape and size vary greatly, their most characteristic property being their colour, and through it alone they may be readily recognised. They may be seen under the microscope to dissolve in caustic potash, and can again be made to crystallise in the rhombic form by neutralisation with hydrochloric acid. The murexide test (see Chapter I.) may be employed in appropriate cases for their detection.

2. Oxalate of Lime.—This substance crystallises in transparent, strongly refracting octahedra (envelope crystals), which are readily soluble in hydrochloric and insoluble in acetic acid (*Fürbringer*).¹⁰⁹

The occurrence of isolated crystals of oxalate of lime (fig. 118) has no clinical significance. They have been found in healthy urine, and their number may be greatly increased after the ingestion of food containing a large proportion of oxalic acid, such as tomatoes, fresh beans, beetroot, asparagus, &c. On the other hand, oxaluria as a morbid state cannot be measured by the use of the microscope alone, since the

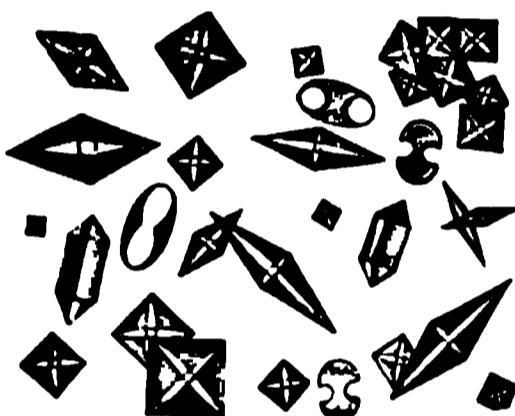


FIG. 118.—Oxalate of Lime from Sediment in a Case of Cystitis and Pyelonephritis (eye-piece III., objective 8A, *Reichert*).

urine may contain a large proportion of oxalic acid, whilst neither it nor its salts are precipitated as crystals. In such cases, the urine must be analysed with a view to ascertaining the proportion of oxalic acid which it contains.

3. Bilirubin and Hæmatoidin.—*Bilirubin* is deposited in the urine either in the amorphous form or in crystals. Crystals of bilirubin have a twofold character, occurring either as clusters of needles, or as minute rhombic tablets, and their colour ranges from yellow to a beautiful ruby-red. They are soluble in caustic soda, and on the application of a drop of nitric acid a green rim forms round them. *Kussmaul*¹¹⁰ has discovered these crystals in jaundice, and *Ebstein*¹¹¹ in pyelonephritis.

Hæmatoidin resembles bilirubin as closely in appearance as in its chemical properties. The crystalline formations of the two are identical (see fig. 95); but it would appear that hæmatoidin may be distinguished chemically from the fact that it turns a transitory blue when treated with nitric acid (*Holm*¹¹²), and is insoluble in caustic potash and æther.

(*Städeler*).¹¹³ According to *Hoppe-Seyler*,¹¹⁴ hæmatoidin and bilirubin are in all respects indistinguishable, and the experience of the author lends support to the view of this great authority; for he has repeatedly had occasion to observe that, in the case of jaundice, the yellow cellular elements, and notably the epithelium, are stained red, passing to blue on the application of nitric acid—a reaction which is assumed to belong only to hæmatoidin, and yet in these cases there can be no doubt that the substance present is bilirubin (see also Chapter I.).

*Leyden*¹¹⁵ found these crystals in nephritis gravidarum, *Foltanek* and *Rosenheim*¹¹⁶ in acute yellow atrophy of the liver, and *Fritz*¹¹⁷ in various chronic and acute diseases, e.g., in a case of carcinoma of the liver, in scarlatina and typhoid. They were usually connected with cellular débris, and in jaundice alone were they also met with in the free state. The author may say that he has repeatedly met with similar forms in severe jaundice of the most distinct types, as in atrophy of the liver,

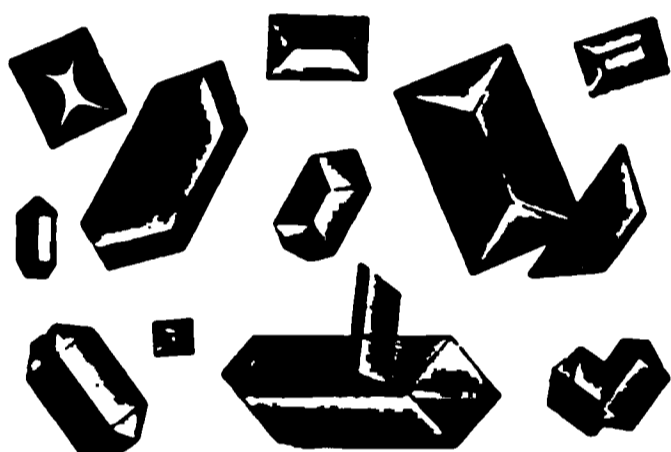


FIG. 119.—Triple Phosphate Crystals from the Sediment in Chlorosis (eye-piece III., objective 8A, *Reichert*).

cirrhosis, and phosphorus poisoning.¹¹⁸ In general it may be said that the presence of such free crystals in considerable quantity implies an antecedent haemorrhage or the bursting of an abscess—as a suppurating echinococcus cyst—into the urinary passages.

4. Triple Phosphates.—Crystals of triple phosphate occur commonly in weakly acid urine, as in the faeces (see Chapter VI.), as bodies of large size and of the coffin-lid form (fig. 119). They are readily soluble in acetic acid. Even when found in large numbers, this fact alone will hardly warrant the diagnosis of phosphaturia.

5. Basic Phosphate of Magnesia.—The crystals of this body have the form of large strongly refracting plates, usually in the shape of elongated rhombic tablets (fig. 120). They are readily soluble in acetic acid, and are precipitated again on the addition of carbonate of soda. They are found in concentrated urine of feebly acid, neutral, or alkaline reaction (*Stein*).¹¹⁹

6. Neutral Phosphate of Lime.—These crystals appear as pointed

wedge-shaped prisms, either singly or in clusters. They are decomposed by ammonia, and dissolve in acetic acid (fig. 121). They occur commonly in feebly acid urine which is becoming alkaline.



FIG. 120.—Basic Phosphate of Magnesia (eye-piece II., objective C, Zeiss).

7. Sulphate of Calcium.—This substance is rarely present in the urinary sediment. It generally takes the form of long colourless



FIG. 121.—Neutral Phosphate of Lime from the Urine of Chronic Nephritis after standing for twenty-four hours. The urine was feebly acid (eye-piece III., objective 8A, Reichert).

needles, but occurs also in elongated tables with abrupt extremities. Amongst such crystals are sometimes to be seen masses of indeter-



FIG. 122.—Calcium Sulphate, Abundant Sediment (eye-piece II., objective C, Zeiss).

minate crystalline structure (fig. 122). They are insoluble in ammonia and acids. Clinically their presence is of little consequence (*Valentiner*,

Fürbringer).¹²⁰ In a patient suffering from a peculiar affection of the ureters and a tendency to concretion in the urinary passages, the urine presented triple phosphates, carbonate of lime, and numerous crystals of sulphate of lime (*v. Jakob*).¹²¹

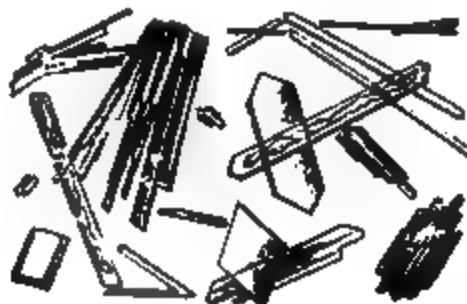


FIG. 123.—Crystals of Hippuric Acid (eye-piece II., objective C, Zeiss).

8. Hippuric Acid.—Crystals of hippuric acid have the form of four-sided prisms, and are scattered separately or in groups (figs. 123 and 124) through the sediment.

They are soluble in ammonia, insoluble in hydrochloric acid. They



FIG. 124.—Hippuric Acid from the Urine of a Rheumatic Patient after large doses of Benzoic Acid had been taken (eye-piece II., objective 8A, Reichert).

occur in considerable quantity after the exhibition of benzoic acid, of the ingestion of certain fruits, as cranberries and bilberries. Otherwise they are a rare manifestation, and have little influence on diagnosis.

9. Cystin.—The crystals of this body are seen as symmetrical hex-

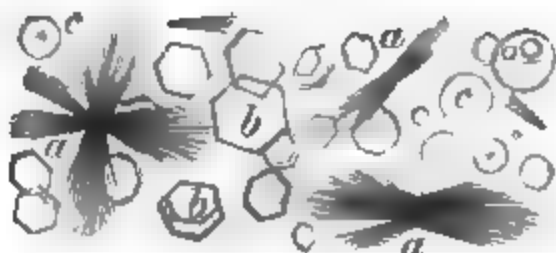


FIG. 125.—a Tyrosin; b. Cystin; c. Leucin (eye-piece II., objective 8A, Reichert).

gonal tables (fig. 125, b), superimposed upon or contiguous to one another. They are insoluble in acetic acid, but readily soluble in ammonia. In this they differ from the crystals of uric acid.

Cystin may also be present in solution in the urine. It may then be precipitated by acetic acid.

When crystals are found in the urine which answer to the description given here, they should be separated by filtration or decanting, the sediment washed with a little water, and the substance in question tested on the platinum foil. Cystin burns with a bluish-green flame, and without melting.¹²²

If cystin be boiled in caustic potash holding oxide of lead in solution, sulphide of lead forms (*Liebig*).¹²³

When heated with caustic potash on a silver plate (as a coin), it leaves a brown or black permanent mark. When dissolved in a boiling solution of caustic potash, dilution with water and the addition of a solution of ferrocyanide of sodium give a violet colour (*J. Müller*).¹²⁴

This reaction, according to *Krukenberg*,¹²⁵ belongs not to the cystin, but to sulphide of calcium, which constitutes an impurity of the caustic potash with which it has been boiled.

10. Xanthin.—*H. Bence Jones*¹²⁶ once found this substance in the urine of a lad who had three years previously exhibited the symptoms of renal colic. It was seen in the sediment in the form of whetstone crystals, which were insoluble in acetic acid and soluble in ammonia (thus distinguished from uric acid). Such bodies are clinically of importance, inasmuch as they may give rise to the formation of calculi (see the observations of *Bence Jones*, l.c.).

11. Tyrosin and Leucin.—These substances generally occur together in the urine.

(a.) *Tyrosin* is seen in the sediment in sheaves of very fine needles (fig. 125, a), which are insoluble in acetic acid, soluble in ammonia and hydrochloric acid.

To determine the character of tyrosin chemically, the sediment containing it should be separated by filtration, washed with water, dissolved in ammonia with the addition of carbonate of ammonia, and the solution allowed to evaporate. The tests for tyrosin may then be applied:—

1. A milligramme of the substance obtained is placed on a watch-glass and moistened with a drop or two of sulphuric acid. The mixture is covered and allowed to stand for half-an-hour. It is then diluted with water, heated, while still hot saturated with calcium carbonate, and filtered. The filtrate is colourless, and when treated with acid-free ferric chloride (compare Chapter IV.) it assumes a violet tint (*Piria, Stüdeler*).¹²⁷

2. When the tyrosin is heated with nitric acid on platinum foil, it assumes an orange-yellow colour. The residue is dark yellow, and turns reddish yellow on the addition of caustic soda. When the latter is evaporated, the substance which remains is of a deep brownish-black colour (*Scherer*).¹²⁸

3. Tyrosin crystals are dissolved in hot water, and the still hot solu-

tion is treated with mercuric nitrate and nitrite of potash. The fluid assumes a dark-red colour, and yields an abundant red precipitate (*R. Hoffmann and L. Meyer*).¹²⁹

4. *C. Wurster*¹³⁰ recommends the following process :—Tyrosin is dissolved in boiling water and a little dried quinone is added. The fluid quickly assumes a deep ruby-red colour, which changes to brown after the lapse of twenty-four hours. The quinone-tyrosin reaction can be depended upon only when tyrosin has been isolated as the free acid [para-hydro-oxyphenol-amido-propionic acid], and will not serve as a test unless it appears readily after the application of heat without prolonged boiling. Quinone alone, or in presence of phenol, gives, when boiled, a rosy-yellow colour to its solutions.

Tyrosin occurs dissolved in the urine as well as in crystals. To obtain it from solution, basic acetate of lead should be added, when a precipitate will form. The fluid should now be filtered, and the filtrate freed from lead by the addition of sulphuretted hydrogen, again filtered, and partially evaporated on the water-bath. The residue is repeatedly extracted with small quantities of strong alcohol, and the extract several times boiled with weak alcohol and then allowed to evaporate spontaneously.

(b.) *Leucin*.—This body, which is commonly associated with tyrosin, is for the most part held in solution in the urine, but to some extent occurs also in the sediment in the form of small spheres (fig. 125, c). The process for its detection is the same as that for tyrosin. It may be separated from the latter by crystallisation from a watery solution, and then purified by recrystallisation from a solution of ammonia in boiling alcohol. When quite pure, leucin crystallises in delicate plates ; when impure, it forms little bulbs of amorphous structure. It may be known by the following tests :—

1. A solution containing leucin, when heated with proto-nitrate of mercury, deposits metallic mercury (*Hofmeister*).¹³¹

2. When consumed with nitric acid on platinum foil, it leaves a colourless residue. This, when heated with caustic potash, forms drops of an oily fluid which does not adhere to the platinum (*Scherer*).¹³²

Tyrosin has been found in the urine in conjunction with leucin in phosphorus poisoning, acute yellow atrophy of the liver, and several of the infectious diseases (*Frerichs, Schultzen, Reiss, Pouillet, A. Fränkel, Blendermann, A. Irsai*).¹³³ *Prus*¹³⁴ has found abundance of leucin in the urine in cases of leukæmia.

It is probable, however, that in some instances where these bodies were supposed to have been in the urine they were not sufficiently identified ; and it is within the author's experience that a deposit has repeatedly been taken as consisting of tyrosin, until subsequent

chemical analysis exposed the error. He is convinced that tyrosin is very rarely present in the urine of phosphorus poisoning.¹²⁵

12. Soaps of Lime and Magnesia.—In the urine of various diseases are often found crystals bearing a very close resemblance in form to those of tyrosin (*r. Jakob*), but possessing distinctive characters of their own. The accompanying illustration (fig. 126) represents such crystals, which the author had the opportunity of examining once only in tolerable abundance. They occurred in the sediment from the feebly acid urine of a woman with severe puerperal septicæmia. They are obviously very similar to those of tyrosin, but yielded none of the characteristic reactions of that body as given above (1-3). The material was not sufficient for further analysis; but it appeared from the behaviour of the substance in question with regard to solubility, &c. (see

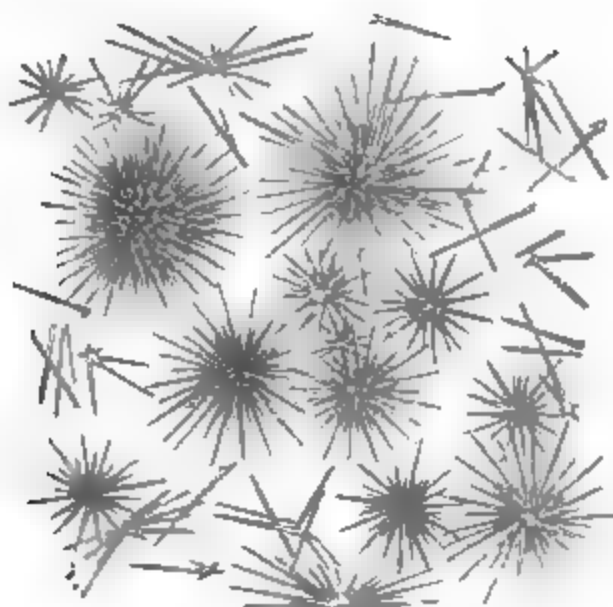


FIG. 126.—Lime and Magnesia Soaps from the Urinary Sediment of a Woman with Puerperal Septicæmia (eye-piece 13., objective 8A, Reichert).

p. 234), that it was probably formed of the lime and magnesia salts of the higher fatty acids.

(b.) Amorphous Deposits.

1. Urates.—Amorphous urates have the appearance of fine granules, disposed singly or in masses. They are entirely dissolved by heating or the addition of acids, and the sediment when treated in this way exhibits free uric acid, for the most part in the form of rhombic tablets.

2. Oxalate of Lime.—The envelope crystals of oxalate of lime have been already described (see Sediments of Acid Urine). This substance appears also as dumb-bell-shaped figures. They are unaffected by acetic acid, and dissolve in concentrated solution of hydrochloric acid.¹²⁶ [The oxalate of lime dumb-bell is really a disc with a central depression on either face, and it presents this appearance when seen sideways (*F.*

Taylor).¹³⁷ Such formations result from slow precipitation in presence of colloid matter (*Orl*).¹³⁸

3. Sulphate of Calcium.—In addition to its crystalline form (fig. 122), sulphate of calcium in the urine takes the appearance of dumb-bell-shaped amorphous masses, which are insoluble in ammonia and in concentrated solutions of hydrochloric acid.

When the sediment contains this substance in considerable quantity, it may be separated from the other constituents by decanting, filtering, and washing in cold water. It is then dissolved in a large bulk of hot water. If to one portion of this solution a quantity of chloride of barium be added, a precipitate of barium sulphate forms, and this is insoluble in nitric and in hydrochloric acid. If another portion be treated with ammonium oxalate, a precipitate of oxalate of lime falls. This precipitate is insoluble in acetic and soluble in hydrochloric and nitric acids.

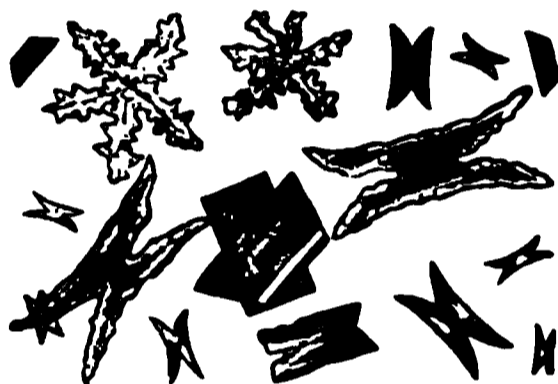


FIG. 127.—Triple Phosphate Crystals (rare form) from Urine in Alkaline Fermentation (eye-piece II., objective 8A, *Reichert*).

4. Brown and Yellow Concretions.—The urinary sediment may contain such concretions either free or associated with cells, and consisting of haematoxin or of bilirubin, which, as we have seen, is perhaps indistinguishable from haematoxin. According to *Holm*,¹³⁹ when a substance of this sort is found to be soluble in caustic potash, and exhibits a coloured ring, of which green forms one zone, on the application of nitric acid, we are to assume that it consists of bilirubin. When, on the other hand, it is insoluble in caustic potash, and colours transitorily blue with nitric acid, he would suppose that it is formed of haematoxin.

5. Fat.—Fat is deposited in the form of strongly refracting globules of varying size, which are readily soluble in æther. It may be present in the urine in small quantities after the fracture of bones, and in chronic inflammation of the kidney, attended with much fatty degeneration of that organ. It occurs in greater abundance, however, only in chyluria, which for the most part depends upon the action of certain worm parasites (*Distoma haematoxylum* and *Filaria sanguinis hominis*), and in phosphorus poisoning. We shall have occasion later to revert to the subject of chyluria and lipuria.

B. Sediments from Alkaline Urine.**(a.) Crystalline Deposits.**

1. Triple Phosphate.—The crystals of triple phosphate deposited in alkaline urine exhibit a very great diversity of character, especially when newly formed in the process of ammoniacal fermentation. They may be seen under such circumstances to resemble snowflakes, or again as peculiar jagged figures, resembling flags or elder leaves (figs. 119 and 127). Their most permanent and characteristic form, however, is that of large colourless and more or less perfect coffin lids or knife-rest shapes.

2. Indigo.—Indigo occurs in the urine as concretions and amorphous fragments, and also in the form of blue crystals and fine blue needles, which mostly cohere in clusters. Crystals of indigo are no very rare manifestation in decomposing and fermenting urine. They are derived from the decomposition of indoxyl sulphate. These crystals were found to be present in remarkable abundance during the process of



FIG. 128.—Indigo Crystals from a Urine rich in Indican, after standing for eight days at ordinary temperature (eye-piece III, objective F, Zeiss).

ammoniacal fermentation in the urine of jaundice, from a patient with hypertrophic cirrhosis of the liver. The urinary sediment in a case of abscess of the liver was recently examined by the author. It contained numerous indigo crystals, and had an *acid* reaction (fig. 128) (*v. Jahsch.*). In another case,¹⁴ one of tabes with cystitis, the urine was so rich in indigo as to derive from the crystals a light-blue colour. The accompanying figure (fig. 128) is partly composed from the preparations of this sediment.

[*Dr. Ord*¹¹¹ has observed a deposit of indigo in the alkaline urine of a patient who had long suffered from enlarged prostate. In this case, besides pure indigo, there were several bodies stained by that substance. Among them were crystals of urates and phosphates, epithelium, yeast-cells, and bacteria. In this way, as *Ord* points out, the form of the indigo is determined by the form of the bodies on which it is deposited.]

3. Urate of Ammonium.—Urate of ammonium forms spherical bodies,

dark in colour and of varying size. Their circumference is beset with radiating spicules of crystalline structure (fig. 129). [Hence they are known as hedgehog crystals.] They are soluble in hydrochloric and acetic acids, with the formation of uric acid, which is deposited in the form of rhombic tables.

4. Phosphate of Magnesia.—The crystals of this salt have been already (fig. 120) sufficiently described.



FIG. 129.—Crystals of Urate of Ammonium, Sediment in Alkaline Fermentation (eye-piece II., objective 8A, Reichert).

5. Cholesterin.—Crystals of cholesterin are very rarely to be seen in the urinary sediment. The author has observed them but once, and that was in the case of a man who suffered from tabes and cystitis. The cholesterin took forty-eight hours to deposit in crystals. The urine when freshly voided had a slightly acid reaction, was turbid, and when shaken, a great number of scaly particles could be seen in it with the naked eye (fig. 130). *A. Glinski*¹⁴² quotes a similar observation.

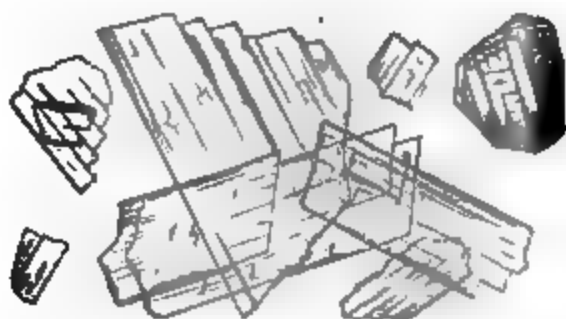


FIG. 130.—Cholesterin Crystals from a Sediment in a case of Tabes and Cystitis; crystallised from ether and alcohol (eye-piece II., objective 8A, Reichert).

(b.) Amorphous Deposits.

1. The large, dark-coloured, spherical bodies represented in fig. 129 consist of *urate of ammonia*. They are distinguished by their solubility in acetic and phosphoric acids, with the simultaneous deposit of rhombic tables of uric acid.

2. Larger or smaller particles, which dissolve in acetic acid without the evolution of gas: *basic phosphatic earths*.

3. Particles of varying size, which are soluble in acetic acid with the evolution of gas: *carbonates of the alkaline earths*.

4. Dumb-bell-shaped masses and coarsely granular concretions, dissolving in acetic acid with the evolution of gas: *carbonate of lime* (fig. 131).

5. *Indigo.* (See fig. 128.)

III. Urinary Concretions.

Concretions of considerable size are occasionally to be seen with the naked eye in the urine (urinary sand). They consist for the most part of urates, or of urates and uric acid together. Their recognition is of great consequence in the diagnosis of renal colic (nephrolithiasis). These concretions are usually more or less stained, and their constitution may be readily determined by the tests already given for the compounds of uric acid. To facilitate their detection the concretion is reduced to powder and the murexide test applied.

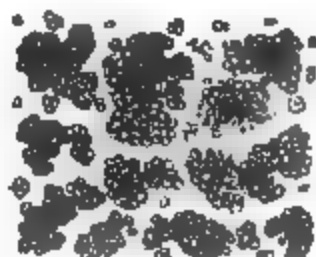


FIG. 131.—Deposit of Carbonate of Lime, Sediment of Ammoniacal Urine (eye-piece II, objective C, Zeiss).

Phosphatic concretions of larger size occur more rarely. They are light-coloured and of little consistence. To determine their nature the concretion in this case also is pulverised and tested for phosphates.

Other concretions occasionally to be met with in the urine are those of cystin, xanthin, oxalic acid, and indigo (*Ord, H. Chiari*).¹⁴³ Those of the last-named substance are easily recognisable by their colour. The chemical tests for other forms of stone, as the oxalate, cystin and xanthin concretions, are treated of elsewhere. (See this chapter: 9. *Cystin*; 10. *Xanthin*; 2. *Oxalate of Lime*.)

In a case which occurred in the author's clinic, and which proved fatal from uremia, the right kidney was found to be cystic, and it contained a large quantity of crystalline concretions. These were ascertained by the author's colleague, Hofmeister, to consist of oxalate of lime, an insoluble proteid, and a derivative of blood pigment.

The constitution of these bodies is dealt with at greater length in the ordinary text-books on the chemistry of urine (see *Huppert*, *Hoppe-Seyler*, and *Leube-Salkowski*).

IV. Cylindrical Bodies visible to the Naked Eye.

1. **Spiral Bodies.**—*R. v. Jakesch*¹⁴⁴ has described bodies resembling *Curschmann's* spirals in the urine of renal stone. They were visible to the naked eye, and consisted of mucin and fibrin. The author has termed the condition *ureteritis membranacea*. A similar appearance is recorded by *Baumüller*.¹⁴⁵

2. **Fibrin Coagula.**—Large and much-matted fibrin clots have been observed by the author¹⁴⁶ in a case of renal abscess probably due to echinococci (fig. 132); and similar observations have been published by *Klein*.¹⁴⁷

It may be mentioned here that *Materba*, *Senna-Saleris*, and *Reale*¹⁴⁸ have noticed that the urine occasionally has a stringy quality (*glischuria*) which they attribute to a special fungus—the *glischro-bacterium*.

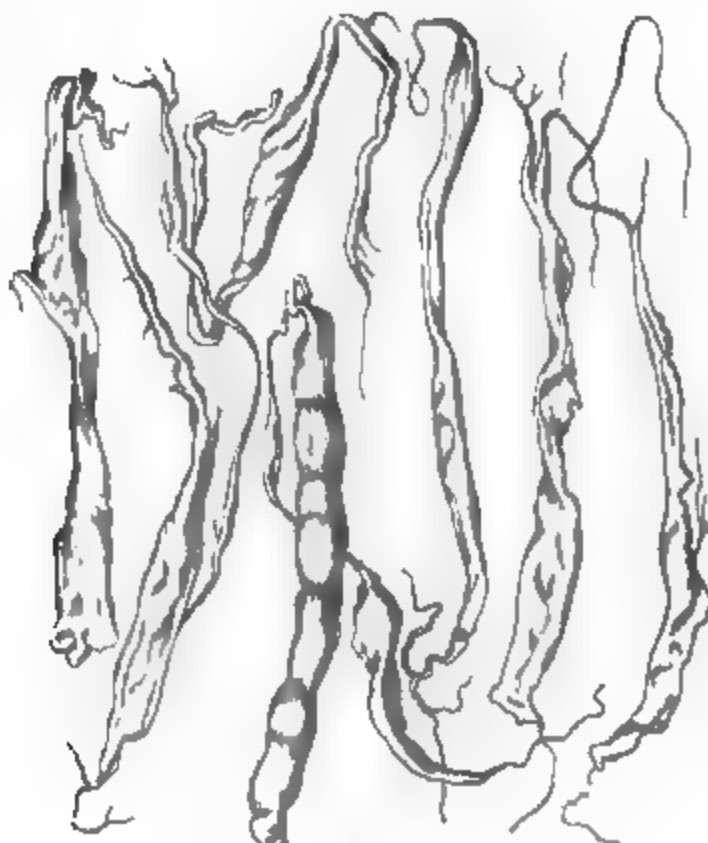


FIG. 132.—Fibrin Coagula from the Urine.

V. Foreign Bodies in the Urine.

As occasional impurities of the urine may be enumerated fatty particles (often introduced with the catheter), fibres of silk, linen, and wool, particles of feathers and wood, and starch granules, the latter being employed as starch powder in medication of the urethra.

The appearance in the urine of substances belonging to the faeces is a fact of serious import. In drawing a conclusion from their presence, care, of course, must be taken to ascertain that they were actually passed with the urine; and when this is so, they are evidence of an abnormal communication between the urinary passages and the gut.

Fragments derived from tumours, as sarcoma and carcinoma, may find their way into the urine by invasion from neighbouring organs.

Hairs have been known to be passed with this fluid (*pilimictio*). They have generally been derived from dermoid cysts in the urinary passages. Sometimes they have been conveyed by accident or design (*hysteria*) into the recipient vessel.¹⁴⁹

III. CHEMICAL EXAMINATION OF THE URINE.

A. Organic Substances.

1. Proteids.—We shall begin with the consideration of albumin, which is the commonest of all the morbid constituents of the urine.

It is still an open question whether albumin ever occurs in considerable quantity in the urine of health. For whilst, on the one hand, the older authorities (as *Frerichs*, *Vogel*, and *Ultzmann*¹⁵⁰) have recorded its occasional presence in healthy urine, and the more recent investigations of *Leube*, *Fürbringer*, *Senator*, and *C. Posner*¹⁵¹ would seem to have established the possibility of a physiological albuminuria, the researches of *v. Noorden*¹⁵² are not less decisive in the opposite direction; and *Leube* and *Winternitz*¹⁵³ from recent very careful experiments conclude that not *every* urine contains albumin. [*Grainger Stewart*¹⁵⁴ has found that albumin is occasionally present in very minute quantity in the urine of persons apparently healthy; and he believes that in some cases it is derived from epithelial and other *débris* after the urine has been secreted at the kidney.]

For the purpose of such investigations the following plan may be adopted from *Leube*:—Normal urine is taken, and it is first ascertained to be free from bacteria and from albumin, as shown by the ordinary methods. It is next distilled in vacuo at a low temperature* (37° to 39° C.), and the residue, with the included sediment, is either tested directly, in the manner to be indicated at p. 297, or it is first treated with alcohol; and of the sediment which remains when the spirit has been driven off, one part is dissolved in water, another in acetic acid, and a third in caustic potash, and the solutions submitted to the tests described at p. 297. Proceeding in this way, it will be found that *Leube's* statement concerning healthy urine is entirely correct, and not a trace of albumin can be shown in it. In the urine of certain diseases, on the other hand, as in valvular heart-disease with compensating hypertrophy, where the most sensitive tests directly applied will fail to evince the presence of albumin, it can readily be shown in the concentrated fluid obtained as above.

Recent investigations, and especially those of *Ott*,¹⁵⁵ into the subject of nucleo-albuminuria, have shown that the albumin present in healthy urine is nucleo-albumin, and, inasmuch as every urine holds this substance, albumin is in this sense a normal constituent. This question

* The distillation in vacuo may be effected by connecting an air-pump with the flask. The procedure recommended by *Anschütz*¹⁵⁵ may be adopted with advantage.

has lately acquired fresh interest from the admirable observations of *Mörner*,¹⁵⁷ who found that chondroitin sulphate and nucleic acid—the latter in very minute traces—were constantly present in urine. He has shown further that normal urine contains serum-albumin, which, on the addition of acetic acid, combines with the substances just mentioned, and that the newly formed compound is identical with the nucleo-albumin of other writers (*Huppert*).

It may at all events be confidently stated that the urine occasionally holds a variable quantity of albumin (*serum-albumin, globulin*) as a temporary constituent, whilst at the same time the kidneys exhibit no alteration of structure, and its presence in such cases is to be attributed to a sudden disturbance of the circulation (see *Schreiber's* experimental albuminuria). To the same category of causes must be ascribed the conditions similarly observed by *O. v. Petersen* in school-children and healthy persons;¹⁵⁸ by *Stirling* in apparently healthy boys; and by *Ringstedt, Hubner, Washburn, Teives, Finot, Capitau, and Beckmann*. [*Pary's*¹⁵⁹ "cyclical albuminuria" occurs during the day and disappears during the night. It seems to be caused by the erect posture. Such a condition is said by *Trissier* to occur especially in young male subjects the offspring of gouty or arthritic parents. *Monon's*¹⁶⁰ remittent albuminuria has probably the same significance. This remittent albuminuria is attributed by *Ribbert* to a reversion on the part of the kidney to the embryonic type, or to its inability to adapt itself after birth. *Fagge*¹⁶¹ has described a paroxysmal albuminuria which he believed to be a form of haemoglobinuria, where globulin resulted from the decomposition of haemoglobin. The condition is occasionally associated with the deposition of oxalates. Finally, a temporary albuminuria may depend upon diet or be induced by exercise (*Taylor*).^{162]} Albuminuria of the like cause and import sometimes occurs, as *Falkenheim*¹⁶³ has remarked, in connection with morbid states. It is important to bear in mind that such cyclical or intermittent albuminuria is sometimes the accompaniment of, and depends upon, chronic inflammatory states of the kidney—such, for instance, was the underlying condition in the case published from the author's clinic by *Ott*,¹⁶⁴ and in another recorded by *Osswald*.¹⁶⁵ [The abuse of morphia is recognised by *Huchard*¹⁶⁶ and by *Haig*¹⁶⁷ as a cause of albuminuria, dependent probably on vascular disturbance.] *Virchow* originally, and after him many other observers, have shown that the urine of newly born infants often contains albumin.¹⁶⁸ This, according to *Flensburg*,¹⁶⁹ is in the form of nucleo-albumin.

*Bright*¹⁷⁰ in England was the first to determine the connection between renal disease, dropsy, and albuminous urine; and after him the labours of such men as *Christison* and *Rayer, Frerichs* and *Traube*,¹⁷¹ formulated what was known of albuminuria as a clinical

symptom. For a long time it was thought sufficient merely to determine the presence of proteid matter in the urine, and the further question was not raised as to whether it occurred in more forms than one. It is now established on clinical and physiological grounds that, in addition to serum-albumin, the urine may contain globulin, peptone, albumose, oxy-hæmoglobin, nucleo-albumin, and fibrin; and it is obviously important to be able to distinguish between these bodies. At present the chief clinical interest attaches to serum-albumin, peptone, and albumose, for by methods now well established we can readily separate each of these from the others. The very simple methods which *Kauder* and *Pohl*¹⁷² have recently communicated for the detection of globulin in serous fluids and urine have placed us in the same favourable position with reference to that body, and by their aid we may hope soon to learn whether globulinuria also exists as an independent condition. In view of their importance in this connection, we shall describe these methods with the others. It follows from what has been said that albuminuria is of different kinds, and we may distinguish: (1) Serum-albuminuria, which alone will be understood to be meant by the term "albuminuria" in the following pages; (2) Peptonuria; (3) Albumosuria; (4) Globulinuria, which as yet is not known to occur by itself;¹⁷³ (5) Fibrinuria; (6) Hæmaturia; Hæmoglobinuria; and (8) Nucleo-albuminuria (Mucinuria).

1. **Albuminuria.**—Under this heading we shall group those states in which the urine is found to contain serum-albumin, in combination, perhaps, with a variable proportion of globulin.

The author has come to the conclusion, as the result of repeated analysis, that the urine of serum-albuminuria does not invariably contain globulin.

Serum-albumin in notable quantity is never found in healthy urine. Its appearance is in all cases a morbid symptom of great importance.

The albumin of urine may be derived from the kidney (renal albuminuria), or by admixture from parts in the urinary passages beyond the kidney (contingent, accidental albuminuria).

(a.) **Renal Albuminuria.**—This form of albuminuria, which is at once the commoner and the more serious, depends in all cases upon disturbance of the renal functions, and such disturbance may be due to various causes. By far the most frequent of these are inflammatory and degenerative changes in the structure of the kidneys. It should be mentioned here that where such processes in the kidney are the underlying cause of albuminuria, we cannot always infer their extent and severity from the quantity of albumin eliminated. So far indeed is this from being the case, that there are certain forms of renal disease of an especially serious character (granular, red, or contracted kidney), in which the urine contains only traces of albumin. Another cause of

albuminuria is found in disturbances of the circulation of different kinds, provided only that such disturbance extends to the vessels of the kidney. It must, however, be borne in mind that errors of the circulation, when of long continuance, may effect changes in the structure (congestion) of the kidney.

To disturbances of the circulation should perhaps be referred the temporary albuminuria of epileptic paroxysms (*M. Huppert*¹⁷⁴), and that which *Schreiber*¹⁷⁵ induced experimentally by compression of the thorax in healthy persons. Possibly, also, that form of albuminuria which often sets in in acute enteritis should be mentioned here (*Singer, Kobler*).¹⁷⁶ As a permanent condition referable to renal congestion may be mentioned the albuminuria of emphysema, heart-disease, weakened heart, &c.

To a third class of causes is to be ascribed the albuminuria of fever (*Leyden*).¹⁷⁷ The circumstances which promote the elimination of albumin in the febrile state are manifold. In the first place, the accompanying changes in blood-pressure would be alone sufficient to account for albuminuria. Then it is known that a long continuance of the febrile processes is competent to induce structural alterations in the renal epithelium, which in their turn lead to albuminuria. It is conceivable, again, that the specific cause (fungi) in certain fevers exerts an immediate influence in the matter, since we find that in many infectious diseases the proliferating micro-organisms are eliminated in large quantities by the kidneys. The admirable experiments recently undertaken by *H. Lorenz*¹⁷⁸ lend great support to the theory that the albuminuria of fever depends directly upon certain histological changes in the renal epithelium, especially affecting the "rodded" lining cells.

There is a fourth form of albuminuria to be distinguished in respect of causation from those already mentioned. It occurs in anaemic and enfeebled individuals, independently of kidney-disease, disordered circulation, and the febrile state; and can be accounted for only by assuming changes in the constitution of the blood, of such a character that, whilst the structure of the kidneys remains intact, and the blood-pressure is not appreciably altered, albumin is allowed to exude with the urine (*v. Bamberger's hæmatogenous albuminuria*).¹⁷⁹

It still remains to say a few words regarding that variety of albuminuria which is intermittent in character. In the author's experience it occurs under the most varying circumstances, and may be either renal or accidental (*q.v.*) in its origin. A number of observations of the kind recently published by *Bull, Mareau, Klemperer, Canfield, and G. Johnson*¹⁸⁰ have proved its occurrence in health, where, no doubt, it is due partly to the circulatory disturbances already referred to, and partly to an inflammatory change in the kidneys.

In the course of a chronic or of an acute nephritis, it will often happen that albumin is found to be temporarily present in the urine (*R. v. Jaksch*).¹⁸¹ When this is so, a careful microscopical examination of the fluid, at a time when it is *free from albumin*, will usually disclose the presence of formed material (casts and renal epithelium) by which the existence of a nephritis may be known. Such intermittent albuminuria belongs especially to contracted kidney. In a case of this disease, if the quantity of urine passed in the twenty-four hours be collected together and examined, it will nearly always be found to contain albumin; but if the same urine be examined in separate portions at shorter intervals—say of one or two hours—it will be found comparatively often that that which is collected in the forenoon is devoid of albumin. But it is not in connection with kidney-disease that intermittent albuminuria is most apt to occur. It is more often associated with affections of the ureters and urethra, and especially with chronic inflammation of the prostatic portion of the latter. In this connection the urine first passed in the morning is generally turbid, and contains albumin, derived no doubt from the pus-cells in the fluid.¹⁸² *Falkenheim*¹⁸³ has recorded a remarkable case of intermittent albuminuria, caused by the pressure of a tumour on the left kidney. In *Pavy's* disease it is probable that a careful microscopical examination would show evidence of an underlying nephritis (see *Merley*).¹⁸⁴ From what has been said above of the various forms of renal albuminuria, it will be seen that this condition is in itself a symptom of very ambiguous significance. Taken with the other physical and microscopical appearances of the urine—but only in conjunction with these—it will afford a valuable indication of kidney-disease. It must be borne in mind that *we are never warranted in inferring the existence of a renal affection or of a nephritis from the mere fact that the urine contains albumin.* This was the error of former times.

(b.) **Accidental Albuminuria.**—The appearance in the urine of albumin derived from a source other than the kidneys is a symptom of much slighter consequence. It may come from the renal pelvis, the ureters, the bladder, or the urethra, or through an abnormal communication from neighbouring parts, *e.g.*, the lymphatics or thoracic duct. In general, a positive judgment as to its source may be formed from the result of chemical and microscopical investigation. Thus, for instance, if the urine exhibits but little serum-albumin and a considerable proportion of pus-cells, the inference is that the former is obtained from the migration of leucocytes in some part of the urinary passages just mentioned. The absence of renal casts and epithelium, moreover, goes far to exclude the possibility of renal albuminuria.

Determination of Albumin (Serum-Albumin).

(a.) **Qualitative Tests.**—The tests at our disposal for the recognition

of albumin are very numerous. The reactions which we shall describe here are all more or less to be depended upon; but there are some amongst them which will engage a larger share of our attention, and they are those which years of clinical experience have shown to be especially trustworthy. Moreover, when applied in the order in which they are given, they will afford a tolerably accurate means of discriminating the various forms in which albumin occurs in the urine.

1. *Nitric Acid and Heat Test*.—A portion of the urine is boiled, and a small quantity ($\frac{1}{10}$ to $\frac{1}{20}$ its volume) of nitric acid (sp. gr. 1.18) added to it. Should a precipitate form on boiling, this may consist either of albumin or of phosphates. If it dissolves on the addition of acid, it is composed of phosphates; if not dissolved in presence of the acid, and if increased thereby, it is albumin (*acid-albumin*).

The application of this test, however, is open to certain fallacies. In the first place, if there be but little albumin present, the quantity of acid usually added will be relatively in excess; and under these circumstances the albumin may combine with the acid to form a soluble nitrate when no precipitate remains. Where, on the other hand, both phosphates and albumin are present, if the acid employed be too little, a portion only of the basic phosphates may be changed into the corresponding acid salts, while the albumin enters into combination with the rest, and remains in solution as an albuminate (union of albumin with a base).

Another occasional source of error is the formation of a precipitate of uric acid. Such a precipitate, however, can usually be known by its deep-brown colour, and from the fact that it is never flocculent. Moreover, it does not fall until the specimen begins to cool.

Finally, the test may be misleading in cases where the urine contains a considerable quantity of resinous acids (pine acids), as happens, for instance, after the use of copaiba balsam. These acids (pinic and pimamic acids) are then precipitated by heat, but can be readily distinguished from albumin by their solubility in alcohol. According to the observations of C. Alexander,¹⁸⁵ this property is useless for differentiating uric acid from albumin, since, under certain conditions, acid albuminate is also soluble in alcohol.

The heat and nitric acid test will serve for the detection of *serum-albumin*, *globulin*, and, where the precipitate falls on cooling, *albumose*, but not for that of peptone.

2. *Acetic Acid and Ferrocyanide of Potassium Test*.—The urine is filtered, and to the clear filtrate a large quantity of acetic acid (sp. gr. 1.064) and a few drops of a 10 per cent. solution of ferrocyanide of potassium are added. If albumin (serum-albumin) be present in considerable quantity, a flocculent precipitate forms; when merely a trace, the fluid becomes turbid or slightly opalescent.

In the latter case it may be necessary to compare it with the filtered urine, in order to estimate the change in its appearance. It will also happen, especially when it contains micro-organisms, that the fluid cannot be rendered clear even by repeated filtration ; and here again the comparison will lie between the simply filtered fluid and that to which the reagents have been added. In the first instance a minimal turbidity, in the second an increase, shows the presence of albumin.

This is a most satisfactory test, and by its means very minute quantities of albumin may be detected. The best and most accurate results, however, may be obtained from its application in the following manner :—A mixture containing a few cubic centimetres of fairly concentrated solution of acetic acid and a little ferrocyanide of potassium is freshly made in a test-glass, and carefully poured upon the surface of the clear filtered urine in a test-tube. If the merest traces of albumin be present, a white ring forms at the point where the two fluids are in contact. For ferrocyanide of potassium, platinocyanide of potassium may be substituted. The test with this reagent, however, is less accurate.

*Jolles*¹⁸⁶ has lately introduced a reagent composed as follows : Perchloride of mercury, 10 grms. ; succinic acid, 20 grms. ; sodium chloride, 10 grms. ; distilled water, 500 grms. The urine is filtered, 4-5 cc. of it are treated with 1 cc. of a 30 per cent. acetic acid, 4-5 cc. of the reagent are added, and the mixture shaken up. The reagent is a substitute for ferrocyanide of potassium, and the test is more efficient.

By this method the presence of *serum-albumin*, *globulin*, and *albumose*, but not of peptone, can be ascertained.

3. *Biuret Test.*¹⁸⁷—The urine is treated with caustic potash, and a dilute (10 per cent.) solution of sulphate of copper is added, drop by drop, with a pipette. If albumin be present, the resulting peroxide of copper (a green precipitate) is dissolved, and the fluid assumes a reddish-violet colour.

This test serves for *albumin*, *albumose*, *globulin*, and *peptone*.

4. *Heller's Test.*¹⁸⁸—The urine is poured carefully, so as to form a layer on the surface of some nitric acid in a test-tube. At the junction of the fluids a white cloud forms in the shape of a ring if albumin be present. This test is very accurate ; but it is not to be recommended for general use, because, in the case of undiluted urine, the deposition of uric acid is apt to cause a brown discolouration, which may easily be mistaken by the inexperienced for the clouding of albumin. After the use of copaiba balsam, too, a similar ring may form.

This reaction is also the basis of an admirable method for approximately estimating the proportion of albumin in the urine (see p. 301).

There are a number of other tests for albumin of a more or less accurate and practicable character, and to some of these allusion must be briefly made.

*Heynsius' Test.*¹⁸⁹—The urine is to be rendered strongly acid with acetic acid;

a few cubic centimetres of a saturated solution of chloride of sodium are added, and the mixture boiled. The presence of albumin is shown by the formation of a flocculent precipitate. Very small quantities can be detected in this way.

[*Potassium Mercuric Iodide Test (Tauret's Reagent)*.—This was found by a committee of the Clinical Society¹⁹⁰ to be the most delicate of a series of reagents examined by it. Tauret's reagent has the following constitution.—Mercuric chloride, 1.35 grms.; potassium iodide, 3.32 grms.; acetic acid, 20 cc.; water, 64 cc. The reaction may also be obtained by *Heller's* method. Serum-albumin, peptone, albumose, alkaloids, and bile acids are precipitated by the reagent. It may be conveniently employed by means of test-papers.]

Hindenlang's (Metaphosphoric Acid) Test.¹⁹¹ If to urine which contains albumin a little solid metaphosphoric acid be added, a precipitate or turbidity forms. This test is very convenient, but will not suffice where only a trace of albumin occurs. The author has repeatedly failed to obtain with it any indication of this substance, while the application of the acetic acid and ferrocyanide of potassium method exhibited its presence. On the other hand, he can confirm the statements of *Penzoldt* and *J. Noorden*,¹⁹² who maintain that a precipitate can often be obtained by this reagent with urines with which all other albumin tests yield negative results.

*Furbringer's Method*¹⁹³ of testing with chloride of mercury and sodium has been shown to be very convenient, especially when the reagent is used in the form of capsules (*Stat's* capsules); but it has no other advantage over those previously described. The use of test-papers also (*as trisures*) in examining for albumin, which has of recent years come into vogue in England and on the Continent, does not seem warranted by experience.

The Picric Acid Test of *Sir George Johnson*¹⁹⁴ is sensitive, but it is not altogether accurate, since this reagent will effect the precipitation of alkaloids and kreatinin (*Jaffe*).¹⁹⁵ in the urine. It must, however, be studied, for it has become the basis of a well known method for the approximate estimation of the quantity of albumin in that fluid. In this connection it will engage our attention later on (p. 304).

Spiegler's Test,¹⁹⁶—The reagent consists of perchloride of mercury, 8 grms.; tartaric acid, 4 grms.; glycerine, 20 grms.; distilled water, 200 grms.

The urine is made acid with a little acetic acid, and poured upon the reagent in a test-tube. At the place of junction of the two fluids a white ring forms when albumin is present. This is perhaps the most delicate test known for albumin, but it is not entirely satisfactory for clinical purposes. Since the reaction occurs, as *Orlitz*¹⁹⁷ has shown, both with nucleo-albumin—a regular constituent of urine—and also with albumose and peptone, the test consequently fails to signify whether the albumin present is pathological, and to discriminate between the different forms of that body.

[Resorcin is used by *Carré* as a test for albumin: 1 grm. is dissolved in 2 cc. of distilled water in a test tube, and the urine is poured on to the surface. The presence of albumin or peptone is shown by a white ring, which, when due to peptone, is dissolved by plunging the test-tube in boiling water.]

Colour Reactions.—Finally, the albumin group yields a number of colour reactions, many of which are applicable to the purpose of detecting its members when contained in the urine. Amongst these are the *Biuret* test already mentioned, and *Millon's* reaction. The *santhoprotein* test, and the colour reactions of *M. Schultze*, *Adamkiewicz*, and *Prokide*,¹⁹⁸ hardly call for special mention here, since their application chemically for the determination of serum albumin has no advantage which does not belong to the methods adopted in the text. For particulars concerning the use of sulphuric and hydrochloric acid as colour-tests for albumin, the reader may consult *Liebermann*, *C. Wierster*, and *E. Salkowski*.¹⁹⁹

Millon's Reaction calls for fuller notice. If to a solution containing albumin mercuric nitrate be added, and heat applied to boiling, the further addition of potassium nitrite will cause the precipitate (if any) and the supernatant fluid to turn red. The reaction is common to all the monohydroxyl-benzol derivatives (*O. Nasse*²⁰⁰). As a test for albumin it is somewhat ambiguous, inasmuch as it can also be obtained from members of the aromatic series. In this connection we shall have occasion to refer to it again.

*Schick*²⁰¹ has applied *Zouchlos's*²⁰² proteid test to clinical purposes. It has no advantage which does not belong also to those described at pp. 297-299. *A. B. Cohen*²⁰³ advocates the use of potassic iodide and the iodide of bismuth and potassium in acid solutions. He claims for this test greater accuracy than it possesses, since alkalies are thrown down by it as well as albumin.

G. Reoch and *J. A. Macwilliam*²⁰⁴ have employed salicyl-sulphonic acid as a means of discriminating serum-albumin from albumoses and peptone. Salicyl-sulphonic acid is readily soluble in water, and if a concentrated solution be added to an acid urine which contains albumin, a turbidity or a precipitate (according to the proportion of albumin present) will result. Should peptone or albumoses be also present, this precipitate diminishes on boiling, and recurs when the mixture cools. An acid reaction is necessary for the application of this test, and should the urine be alkaline, it must be treated previously with acetic acid. In the author's experience, the test is very suitable for distinguishing albumin from albumoses and peptone, as *Macwilliam*²⁰⁵ has pointed out. *r. Engel*, however, contends that it is not in any way to be preferred to the method of *Hofmeister*,²⁰⁶ detailed on a later page.

[*Macwilliam* applies the test in the following way:—About 20 drops of the urine is placed in a very small test-tube, and one or two drops of a saturated watery solution of the reagent is added, or a little more if the urine is strongly alkaline. The tube is rapidly shaken and the fluid at once inspected. Opalescence or turbidity recurring immediately, *i.e.*, in one or two seconds, is an indication of greater delicacy than the cold nitric acid test. If turbidity appears after a longer time, as from one-half to two minutes, this implies the presence of a very minute trace of albumin. The fluid is next boiled. The precipitate, if due to serum-albumin and globulin, does not disappear, but becomes flocculent, while a precipitate from the presence of albumoses and peptone clears up before the boiling-point is reached. For salicyl-sulphonic acid as a test for deutero-albumose and peptone, see p. 311.]

*B. Vas*²⁰⁷ and *A. Ott*²⁰⁸ report most favourably of this as a test for albumin, and the latter strongly recommends it as a bedside expedient. The same authority²⁰⁹ has investigated the value of certain other new tests, as those of *Jolles*, *Tauret*, *Roberts*, *Millard*, and *Raabe*, and finds that some of them are valuable, but without special advantages.

The first three tests described above, when taken together, afford a means of discriminating between the different forms of albumin which are apt to occur in the urine.

(1.) If in a given case the tests 1-3 should all give positive results, it may be concluded that the proteid present is in the form of serum-albumin, usually in conjunction with a small proportion of globulin; but the presence or absence of peptone or albumose in addition remains undetermined.

(2.) If the urine contains but a small quantity of serum-albumin alone, only the first two tests will give any result.

(3.) If the first test fails to disclose the presence of albumin, while a precipitate is formed on the addition of acetic acid in the second, this may consist of nucleo-albumin (mucin) or resinous acids. In the latter case it is soluble in alcohol.

(4.) If, on the application of test 1, no result is at first obtained, but a precipitate forms on cooling the specimen, this precipitate may be removed by filtration and the fluid submitted to test 3 (biuret test), when the characteristic reaction will point to the presence of albumose. In such a case the inference is strengthened if the pure or diluted urine responds to test 2, and if test 3, when applied to the urine directly, produces an abundant precipitate. Further investigation may then be conducted in the manner indicated at p. 305.

(5.) Again, if tests 1 and 2 are unattended with any result, if acetic acid alone will not cause a precipitate, and if test 3 at the same time discloses the presence of albumin, it may be concluded with certainty that the urine contains a large proportion of peptone. This condition is doubtless very rare. The author has met with it several times—repeatedly in the resolution stage of severe pneumonia, in a case of acute rheumatism, when serious and extensive joint symptoms were rapidly disappearing under the influence of preparations of salicylic acid, at some period of phosphorus poisoning (*W. Robitschek*),²¹⁰ and in scurvy (*v. Jaksch*).²¹¹ For the more accurate determination of peptone, it will usually be necessary to resort to the methods described at p. 309.

By proceeding in the order indicated, it will be possible to form some judgment as to the character of the proteids under investigation, and this, as we shall see presently, is often a point of interest clinically.

(β.) Quantitative Estimation of Albumin.

1. *By Weight.*—A certain quantity of the urine—60–100 cc., according as it is rich in albumin or otherwise—is placed in a beaker, and heated on the water-bath. A two per cent. solution of acetic acid is then added, drop by drop, until the albumin has separated in flocculent masses, when the fluid is boiled, and passed through an ash-free filter of known weight. The precipitate on the filter is washed successively with water, alcohol, and æther, and heated to a constant density at 120°–130° C. The filter is again weighed, and the difference in weight due to the precipitate is that of the albumin in the quantity of urine taken. *Deroto's*²¹² plan is still better. The albumin is precipitated with ammonium sulphate, coagulated in the steam-bath, and the precipitate washed with boiling water until the filtrate no longer becomes cloudy on standing, or with the addition of sodium chloride. The

precipitate is then washed with alcohol and æther, and the remainder of the process is conducted as before. To ensure greater accuracy in the result, it will be necessary to ascertain the amount of ash in the filter, and to allow for it. This may be accomplished by burning the filter with the albumin upon it in a platinum crucible of known weight.²¹³ For such determinations the use of glass-wool filters is to be recommended.

2. *The Methods of Roberts, Stolnikow, and Brandberg*²¹⁴ for the approximate estimation of albumin are all based upon the principle of *Heller's* test, and are highly serviceable. The principle referred to may be expressed by the fact, that the richer the urine is in albumin, the more quickly turbidity is developed in the use of Heller's test. If in 100 cc. of urine there is 0.0034 gramme (*Roberts*) or 0.004 gramme (*Stolnikow*) of albumin, the fluid begins to cloud after the lapse of 35–40 seconds, and becomes plainly turbid in 1½ minute.

Brandberg's modification of *Roberts'* and *Stolnikow's* methods gives the best and most accurate results. This observer has found that in a solution of one part albumin in 30,000 of water, and consequently in urine holding 0.0033 per cent. of albumin, Heller's reaction will take place in 2½–3 minutes, and upon this fact he has based his method. It is conducted thus:—The urine to be examined is first submitted directly to Heller's test. If a precipitate is obtained, a measured bulk of the fluid is diluted in a graduated tube with nine times its quantity of water (1 in 10 urine), and Heller's test is again applied to the mixture in the following manner:—In a test-tube of about 1 cm. diameter, some pure nitric acid is placed with a pipette, and in such a manner as not to touch the sides of the tube. The latter is now inclined, and the diluted (1 in 10) urine is carefully poured from a graduated burette along its lower surface, so that it may float upon the acid without blending. Two cc. may be placed in the test-tube in this way. Suppose now that an obvious clouding of the fluid takes place *before* the lapse of three minutes, the diluted (1 in 10) urine contains more than 0.0033 per cent., and the pure urine therefore more than 0.033 per cent. of albumin. If, on the other hand, the albuminous cloudy ring takes longer than three minutes to develop, it is known that the simple urine holds less than 0.033 per cent. albumin in solution. In the first case the mixture is further diluted by *Brandberg* in this manner:—Five test-tubes are taken, and in each he places 2 cc. of the diluted urine. To the first he then adds 4 cc. of water; to the second, 13 cc.; to the third, 28 cc.; to the fourth, 43 cc.; and to the fifth, 58 cc. He then repeats the test with each of these. If in the case of one of them the reaction occurs in the space of 2½–3 minutes, it follows that the fluid in the particular test-tube contains 0.0033 per cent. albumin; and from this the propor-

tion of that substance in the undiluted urine may be directly calculated by the following formula :—

$$p = \frac{k+x}{k \cdot 30}$$

Where

p = percentage of albumin in the undiluted urine.

k = the quantity of 1 in 10 diluted urine in each test-tube.

x = the quantity of water used for dilution.

Brandberg has compiled for the use of the practitioner a practical table by which the percentage of albumin can be readily computed from these data without the trouble of calculation. It is reproduced here with slight modification.²¹⁵ Its conclusions are based upon the supposition that turbidity ensues three minutes after the application of Heller's test :—

I.	2 cc. 1/10 urine,	0.05	0.10	0.15	0.20	0.25	0.30	0.35	0.40	0.45	0.50	% albumin.
II.		1	4	7	10	13	16	19	22	25	28	cc. water.
I.	2 cc. 1/10 urine,	0.55	0.60	0.65	0.70	0.75	0.80	0.85	0.90	0.95	1.00	% albumin.
II.		31	34	37	40	43	46	49	52	55	58	cc. water.
I.	2 cc. 1/10 urine,	1.05	1.10	1.15	1.20	1.25	1.30	1.35	1.40	1.45	1.50	% albumin.
II.		61	64	67	70	73	76	79	82	85	88	cc. water.

The figures in the horizontal line marked II. indicate the quantity of water in cc. which must be added to the 2 cc. of the 1 in 10 diluted urine in order that turbidity may occur in three minutes. In line I. are exhibited the corresponding percentages of albumin in the specimen of urine taken.

In applying this method, it will be convenient to have a burette filled with the 1 in 10 diluted urine, and another containing distilled water. In each of one series of test-tubes may then be placed an equal quantity of nitric acid, with the precautions indicated above ; and in those of another, 2 cc. of the 1 in 10 diluted urine and varying quantities of distilled water (see table). A preliminary experiment is first made, and, according as this experiment shows an excess above or deficiency from the standard percentage of albumin, more or less diluted mixtures of 2 cc. 1/10 urine + water are prepared. These mixtures of urine and water in the second series are now tested by placing them with a pipette on the surface of the nitric acid in the test-tubes of the first series, until that one is reached in which the albuminous ring takes precisely three minutes to form, and from this the proportion of albumin in the urine is calculated. To take an example :—Suppose it is found in this way that turbidity occurs after the lapse of three minutes from the addition to the nitric acid of the fluid in a particular test-tube, and that this fluid has been formed by the admixture of 13 cc. of water to the 2 cc.

of 1 in 10 diluted urine: reference to the table will show that 0.25 corresponds to this number, 13, and consequently the urine contains 0.25 per cent. of albumin. If, however, the fluid in the test-tube to which 13 cc. of water has been added fails, when tested, to become turbid, or does so only after a longer interval, a less highly diluted mixture must be examined in a similar way—as, for instance, the contents of the tube to which 10 cc. water have been added, and so successively with the others (7 and 4 cc. dilution), until the required result is obtained. If, on the other hand, the mixture in the test-tube diluted with 13 cc. exhibits turbidity directly, the test must be repeated with those towards the other extremity of the scale—namely, those containing 16, 19, 22, &c., cc. respectively—until, as before, one is found in which the phenomenon occurs after the lapse of precisely three minutes. Let this be the fluid formed by the admixture of 25 cc. from the burette containing distilled water; then the table will be found to give 0.45 as the corresponding figure, and the urine under examination, therefore, contains 0.45 per cent. albumin.

This method, when carefully conducted, is altogether satisfactory. *Hammarsten*²¹⁶ has compared its results with those obtained by the process of weighing, and has found that the two correspond within 0.266 per cent.

3. *Estimation of albumin by precipitation with picric acid and Esbach's albuminimeter.*²¹⁷

The advantage of this proceeding is its simplicity. It is open to certain fallacies, contingent upon the unsatisfactory character of the reagent used; but it calls for description here as affording the physician a practical, if only approximate, means of determining the proportion of albumin in the urine.

Ten grms. of pure picric acid and 20 grms. of pure citric acid are dissolved in 900 cc. water. When the solution is thoroughly cool, water is further added to 1000 cc., and the mixture is employed as a precipitating agent. An instrument called the albuminimeter is further needed. This is a vessel resembling a test-tube, of stout glass. At its upper part it bears a mark, *R*, and lower down another mark, *U*. The lower third is graduated, and the divisions, which are numbered from $\frac{1}{2}$ -7, are so disposed that the intervals between successive units diminish from below upwards (fig. 133). For the purpose of the experiment, the albuminimeter is used in the following manner:—It is filled with urine up to the mark *U*, and the reagent is then added until the surface of the fluid reaches to the mark *R*. The thumb is then applied to the mouth of the instrument, and it is reversed several times until the fluids are well mixed. It is then covered with an india-rubber stopper, and allowed to stand upright for twenty-four hours. After this the height of the

sediment in the tube is read off by the scale upon its surface. This is constructed so that the numbers express the proportion of albumin in grammes to the litre. Should it happen in a particular case that the upper limit of the sediment overtops the number 7 at the highest point of the scale, i.e., where the proportion of albumin exceeds .7 per cent., the experiment must be repeated with diluted urine. And it will save time if the urine be diluted at the outset, wherever it has been shown by a qualitative test to be comparatively rich in albumin. It occasionally happens that the albuminous precipitate will not cohere, or it may remain floating in the fluid. In such cases the method is inapplicable.

The quantity of albumin present can be estimated only approximately by this method. The experiments of *Czapek*²¹⁵ have shown that its merit is enhanced if certain precautions be observed in its use. The urine to be tested should be fresh and acid, and of low specific gravity, to which end it may be diluted. The fluid should not contain more than four grms. albumin to the litre, and should be kept for twenty-four hours in the albuminimeter at a moderate temperature before the reading is taken. Finally, the figures on Esbach's instrument are too low. The best results are obtained where the urine is below 1.010 specific gravity, and does not hold more than 0.2 per cent. albumin. The conclusions set forth here are confirmed by *Sokolow* and *Th. Geisler*.²¹⁶ *Christensen's*²²⁰ albuminimeter is applied on the same principle, tannic acid being used to throw down albumin. With the instrument the process may be performed more rapidly, but the results are less accurate than with *Esbach's* (*Geisler*,²²¹ *Wavor*, *Federer*).

This method is appropriate to the case of transitory or febrile albuminuria, and it is inapplicable where the urine may be supposed to contain quinine, antipyrin, or thallin.

From comparisons made by *Richter*, it appears that Esbach's method is far less accurate than *Brandberg's*. It is open to very considerable objections, and can be relied upon only within broad limits.

*Noel Paton*²²² employs the albuminimeter to determine the proportion of albumin in a urine. From another portion of the same urine he precipitates globulins with magnesium sulphate, as *Hammarsten* does, and again tests the "globulin free" urine with Esbach's albuminimeter. From the difference in the two results he infers the quantity of globulins present. This plan is commended by its author, but the process obviously partakes of the uncertainty which is inseparable from the use of the albuminimeter.



FIG. 123.
Esbach's Albu-
minimeter.

It may be noticed that *Huppert* and *Záhř*²²³ have endeavoured to found a quantitative method of estimating albumin upon the comparative specific gravity of the urine. Its efficiency has not yet been sufficiently tested, but it may prove to be of service clinically.

2. Peptonuria.—This condition has been invested with very great interest since the discovery by *Hofmeister*²²⁴ of a comparatively simple chemical test for the detection of peptone.

By peptone is here meant *Brücke's* peptone, not the "Kühne" peptone, which hitherto has not been discovered in animals elsewhere than in the contents of the stomach.

From the clinical point of view it is immaterial that the bodies spoken of here as peptones are chemically albumoses. To be consistent in this way, we should substitute for peptonuria the terms deutero-albumosuria, proto-albumosuria, &c.

At the same time, it is necessary, in opposition to other writers (*van Noorden*, *Senator*, *Stadelmann*),²²⁵ to enforce the view that albumosuria thus limited is to be distinguished very sharply from the other or clinical albumosuria, since their clinical significance is widely different.²²⁶

So far as our present knowledge extends, the causes of peptonuria are quite different from those to which the other forms of albuminuria are due. Neither nephritis, circulatory disturbances, nor anaemia will bring about the appearance of peptone in the urine. Its presence there is most commonly—though not invariably—associated with such processes as are characterised by the collection and subsequent destruction of leucocytes under such circumstances that the products of disintegration, including the peptone constituent, of these bodies can obtain admission into the blood-stream, to be subsequently eliminated by the kidneys.

To the condition arising in this way the name *Pyogenic peptonuria* has been given (*Hofmeister*, *Maixner*, *v. Jaksch*).²²⁷ It occurs chiefly in the resolution stage of pneumonia, in purulent pleuritic exudation, and, in general, in suppuration wherever situated, provided that the conditions are favourable to the absorption of the constituents (peptone) of the pus.

Peptone has further been found abundantly in the urine in purulent meningitis, acute articular rheumatism, the suppuration of phthisis—briefly, in nearly all those states which are attended with the formation and breaking down of pus.

Similar results were obtained by *Krehl* and *Mathes*²²⁸ lately by means of another method, notwithstanding that these observers determined the presence only of albumose, while in the cases mentioned *peptonuria* was established.

It follows, therefore, that the detection of peptone in the urine goes far to warrant the inference that suppurative changes are in progress in some part of the system, but such an inference must, in all cases, be strengthened by the exclusion of the other known causes of peptonuria.

Amongst these are cases of scurvy (*inorganic peptonuria*), (*v. Jaksch, Boeri*)²²⁹; and *W. Robitschek*²³⁰ has commented upon the occurrence of peptonuria in phosphorus poisoning. Tissue destruction generally is known to produce the same condition. Such causes as these have therefore to be excluded. *Maixner*²³¹ has also shown that in ulceration of the intestine, peptone derived from the food may pass directly into the blood through the ulcerated parts, and so give rise to this condition (*enterogenic peptonuria*), an observation confirmed indirectly by those of *Pacanowski*.²³² The cases reported by *Chrostek* and *Stromayer*²³³ must also be ascribed to enterogenic peptonuria. *Fischel*²³⁴ has shown that it is normally a constituent of the urine in the puerperal state (*puerperal peptonuria*).

These facts are mentioned chiefly to show that peptonuria does not always imply suppuration; but these other conditions being excluded, it is a valuable diagnostic sign of that process. It is, moreover, available as a means of prognosis and of testing the progress of certain diseases attended with the resolution of pus. Thus, when peptone is found in the urine in the course of pneumonia, it indicates that the stage of resolution has begun. Again, in connection with abdominal tumours or pleuritic effusion, it shows their purulent character; and in purulent meningitis its manifestation varies with the severity of the disease—peptonuria occurring together with a relapse, and so on.

As a means of discriminating between tubercular and epidemic cerebro-spinal meningitis and multiplex haemorrhagic encephalitis,²³⁵ the presence of peptone in the urine is occasionally a fact of crucial significance. It is characteristic of the second disease, occasional in the last, and its absence in presence of the clinical symptoms of meningitis generally implies a tubercular character. Obviously, however, peptonuria may arise accidentally in the course of tubercular meningitis; and in basing a diagnosis upon this condition, care must be taken to ascertain that it is not due to ulcerative processes in other organs, and especially to exclude implication of the lungs.

Again, in the condition which has been called “sepsis occulta,” and which is commonly so difficult to recognise, peptonuria is an important symptom. By its aid especially it will be possible to distinguish the symptoms of septicæmia from those of latent disseminated sarcoma, which present quite a similar clinical character (high fever, rigors).

In a case which came under *Prof. Nothnagel's* care, there had been rigors and a high temperature maintained for a long period, and no other symptoms whatever. These were ascribed to the formation of a deep-seated abscess. The urine was repeatedly examined for peptone without result. The post-mortem showed disseminated sarcomatous nodules.

The author has been engaged for many years in investigating the subject of peptonuria, and he feels warranted in asserting that it

clinical value as a symptom is very great. It may be that further experience will extend and distribute its import ; but there can be little doubt that the significance attaching at present to the form with which we are best acquainted—that of *pyogenic peptonuria*—will not diminish with the progress of knowledge. Recent investigations tend in the fullest manner to confirm this view.²³⁶ Peptonuria is frequently met with in connection with syphilis (*Poehl*).²³⁷ Modern science furnishes an adequate explanation in many instances where the condition is found ; for, granting that micro-organisms have the property of changing albumin into peptone, it is probable that they can also cause the latter to appear in the urine (*Mya, Belfanti, W. Robitschek*).²³⁸

Detection of Peptone.—The methods of *Hofmeister, Devoto, and Salkowski*²³⁹ will be described here.

1. *Hofmeister's Method.*—The urine should first be tested for albumin by the three processes described above (pp. 297–298). If tests 1 and 2 give no result, and no precipitate forms on the addition of acetic acid alone, the presence of peptone may be shown by the biuret test (see p. 298), (*v. Jaksch*),²⁴⁰ but only when this body is in great abundance. If it is not so, the result in this case will be also negative. A further preliminary test may be applied by adding first concentrated acetic acid, and then a mixture of acetic and phospho-tungstic acids. If clouding takes place either directly or after the lapse of a short interval, it may be concluded that peptone is present. The inference is rendered still more certain if, before the application of the test, a little neutral acetate of lead be added (to precipitate mucin) until a flocculent precipitate appears. If the test again gives positive results, peptone is present ; if it remains negative, the urine is peptone-free. This, however, only holds good where the urine contains a considerable amount of peptone.

Hofmeister's test is more accurate. Assuming that the previous methods have failed to disclose albumin, the urine is treated with neutral acetate of lead and filtered. The clear filtrate, which should amount to not less than 500 to 600 cc. in volume, is acidulated with hydrochloric acid, and phospho-tungstic acid is added until a precipitate ceases to form with it. The fluid is then filtered without delay.

Phospho-tungstic acid may be prepared thus :—Commercial tungstate of soda is dissolved in boiling water and phosphoric acid added until the mixture exhibits an acid reaction. It is then allowed to cool, rendered strongly acid with hydrochloric acid, and filtered after standing for twenty-four hours (*Huppert*).²⁴¹

The precipitate consists of peptone combined with phospho-tungstic acid, and various other substances (ptomaines, &c.). It is now washed on the filter with five parts concentrated sulphuric acid in 100 of water, until the fluid which passes through is colourless. In this way the salts

are got rid of. The precipitate, while still wet, is washed from the filter with as little water as possible, and is received in a watch-glass. Barium carbonate is added until the mixture is alkaline, and the latter is then placed on a water-bath at boiling-point, and heated for ten to fifteen minutes, and the biuret test (p. 298) applied. Peptone is shown by the formation of a colour ranging from bluish red to violet, and varying in intensity according to the quantity present. If this be only a trace, the resulting colour is of a dirty red or dull violet. The accompanying precipitate of baryta need not confuse the experiment; but should any doubt exist as to its result, the entire preparation may be placed in a test-tube and allowed to stand for a few minutes. The precipitate then falls to the bottom, and the fluid displays the characteristic tints—from dull red to violet—if it contains peptone. If that body be absent, it has a greenish colour.

If either of the first two tests (p. 297) should show the presence of albumin, whilst the addition of acetic acid and ferrocyanide of potassium after filtration causes only *very slight* turbidity, the albumin must be removed by combination with a metallic oxide—and best with oxide of iron—in the following manner:—A solution, first of acetate of soda and then of perchloride of iron, is added to the urine. This is exactly neutralised with caustic potash, boiled, filtered, and allowed to cool. Tests 1 and 2 are next applied. If neither discloses albumin, and if also with test 2 no blue colour forms, thus showing that the fluid is free from iron, the further process is that described above—hydrochloric acid is added, and a precipitate obtained by the use of phosphotungstic acid, and so on, as before.

If, however, after the precipitation of albumin in the manner indicated, one of the tests alluded to should show that the fluid is not yet free from that body, the previous tests must be repeated until the filtrate exhibits no trace either of albumin or of iron. Should the former substance be present in the urine in great quantity, it may be removed in large proportion by heat, and what remains may be subsequently got rid of in the manner described above.

The application of this method in the case of highly coloured albumin-free urine has the contingent advantage that it involves the removal of colouring matters whose presence might be a source of fallacy. *J. A. Schulter*²⁴² recommends that the urine be first saturated with ammonium sulphate, and the filtrate proceeded with as before. The quantitative estimation of peptone may be effected by the colorimetric process of *Hofmeister* and *Maixner*.²⁴³ Other processes, as that of *Stadelmann*,²⁴⁴ directed to the same end, fail in their purpose, and lead to results which are not trustworthy.

2. *Devoto's Method*.—This test may be applied in the following

manner, and the description differs but little from that given by its author.²⁴⁵

This and *Hofmeister's* test have given identical results when the author has employed them for the investigation of urine. When, however, they were applied to the examination of the blood and the viscera, there was less conformity between them, *Hofmeister's* usually indicating peptone where *Devoto's* failed to do so.²⁴⁶

To 200–300 cc. of urine is added pure crystalline ammonium sulphate in the proportion of 80 grms. to 100 cc., and the fluid is placed in a beaker in a boiling water-bath for half-an-hour, at the end of which the greater part of the salt should have dissolved. It is then steamed for half-an-hour in a *Budenberg's* steam-steriliser, the vapour being kept at 100° C. In this way all the proteids (serum-albumin, globulin, haemoglobin, deutero-albumose, peptone, nucleo-albumin) are precipitated, but only serum-albumin, globulin, and nucleo-albumin (mucin) are thoroughly (and haemoglobin partly) coagulated. The fluid having been heated to 100° C., is at once filtered. The filtrate should be straw-coloured, and free from albumin, as indicated by tests 1 and 2, p. 297. A slight cloudiness appearing quickly with test 2 does not necessarily imply the presence of albumin. A decided turbidity or a precipitate would be due to a proto-albumose, or more probably hetero-albumose. Should the hot filtrate be cloudy, or give the proteid reactions (pp. 297–298), the investigation has miscarried, and must be repeated from the beginning. The residue on the filter is washed first with hot and then with cold water. The resulting filtrates have a more or less decided brownish tint. These are collected, and to one portion of the fluid acetic acid and ferrocyanide of potassium are added to test for albumin. Should no result be obtained, the biuret test is performed with a portion to which caustic soda has been added in excess. Any albumin shown to be present is certainly peptone. The filtrate from the hot washings may exhibit it, but it often happens that peptone first becomes recognisable with the biuret test in the filtrate derived from the cold washings. Several specimens both of the hot and cold washings may be tested until a positive result is obtained.

3. *Salkowski's Method.*²⁴⁷—To 50 cc. of urine are added 5 cc. of hydrochloric acid, and then phospho-tungstic acid till a precipitate ceases to form. The latter is made to agglomerate by cautiously heating, the overlying fluid is drawn off, and the precipitate repeatedly washed with water, dissolved in 0.5 cc. of caustic soda solution (sp. gr. 0.16), heated till the blue or greenish colour disappears, and then tested for the biuret reaction.

This method is expeditious, and therefore eminently practical.²⁴⁸ It has been proved by *E. Robitschek*, in the author's clinic, to yield results

entirely in accord with the other methods given here. The urine to be examined must be free from albumin. If not already so, it should be rendered free by one of the processes described at p. 301 and p. 308. More recent observations have shown the author that, in the case of urines rich in urobilin, the biuret reaction can only be employed with great care, since such urines are capable of giving positive results under this test, even in the absence of peptone.

[*S. Martin*²⁴⁹ has found that in many cases of supposed peptonuria—especially of peptonuria in connection with purulent disease—the morbid constituent of the urine was deutero-albumose and not peptone. The two are distinguished by their behaviour in solutions with ammonium sulphate; deutero-albumose is precipitated by saturation with this salt, while peptone is not.

*Macwilliam*²⁵⁰ recommends salicyl-sulphonic acid as a test for albumoses and peptone. Primary *albumoses* are precipitated by a saturated watery solution of the acid; the precipitate disappears with heat, and reappears when the liquid cools. *Deutero-albumose* is precipitated by the reagent when the fluid examined is mixed with two to three times its bulk of a saturated solution of ammonium sulphate. *Peptone* is readily precipitated by salicyl-sulphonic acid in saturated ammonium-sulphate solutions, and the precipitate clears up on heating, and reappears when the liquid cools. The peptone precipitate is dissolved by the addition of water, glycerine, or excess of the reagent.]

3. Albumosuria.—The distinctive character of this condition was formerly thought to be the presence in the urine of a single body, to which the name of propeptone or hemialbumose had been given. The researches of modern times, however (*Kühne, Chittenden, Herth*²⁵¹), have placed the matter in a new light, and the first two observers have sought to show that propeptone is really a mixture of four different proteids. However this may be, these interesting observations cannot as yet be brought to bear upon our clinical knowledge.

Albumose has been found in the urine in connection with various diseases—as osteomalacia, dermatitis, intestinal ulcer, &c. (*Senator, Ter Gregoriantz, R. v. Jaksch*).²⁵² In severe cases of osteomalacia, which were recently under treatment by *Professor Nothnagel, r. Helly, Schauta*, and himself, the author failed to discover albumosuria; neither has he detected it in advanced rickets. *Raschkr's*²⁵³ records a case of senile osteomalacia in which a supposed albumosuria occurred.

*Loeb*²⁵⁴ believes that he has discovered propeptone in the urine in measles and scarlatina (in this he is confirmed by *Heller*²⁵⁵), and from the observations of *Kahler* and *Huppert*²⁵⁶ albumosuria would seem to be a frequent occurrence in inflammation of the medulla of bones. Later investigations, as those of *Ribbink*,²⁵⁷ support this view. *Köppner*²⁵⁸ has observed albumosuria in mental derangement. The clinical significance of this state is modified by the fact that *Posner*²⁵⁹ has detected propeptone in the seminal fluid (see Chapter IX.). The

presence of albumose, when it occurs alone, may be determined by the consideration of results derived from the tests given for the detection of albumin generally. If, on the application of tests, a precipitate first forms on cooling the specimen, or when it has been allowed to stand for a long time (*vide supra*), and if this precipitate, when separated on the filter, is shown by the biuret test to consist of albumin, a fresh specimen of the urine may be submitted to test 2. In doing this, it may be necessary to add water, since albumose is readily soluble in concentrated salt solutions, and therefore in concentrated urine. Whether with or without this precaution, a further precipitate with test 2 suggests the presence of albumose. Another specimen should now be saturated with common salt, and further treated with acetic acid. If albumose be present, a precipitate forms, which, after the addition of a large amount of acetic acid, dissolves when heated and reappears on cooling. A nitrogenous body resembling albumose was found by *Thormählen*²⁶⁰ in the urine in a case of hydatids of the liver, with jaundice and nephritis.

When the urine holds albumose in conjunction with serum-albumin, the latter must be removed by boiling with acetic acid and chloride of sodium before the tests are applied.²⁶¹

4. Globulinuria.—Globulin probably never, or almost never, occurs alone in the urine, but generally in conjunction with serum-albumin. Consequently its import in disease is the same as that of the latter body. [Serum-albumin is usually present in greater quantity, but in severe organic renal disease and in diabetes *Maguire*²⁶² finds that the proportion of globulin to albumin is often 2.5 : 1. *Senator*²⁶³ asserts that more globulin is present in lardaceous kidney than in other forms of Bright's disease.]

To *Kauder*²⁶⁴ belongs the credit of discovering a simple method for the detection of this body in presence of serum-albumin. The urine is rendered alkaline with ammonia, allowed to stand for an hour, then filtered, and to the filtrate is added its own volume of a saturated solution of sulphate of ammonium. If globulin be present in any quantity, a flocculent precipitate falls.

By an extension of this method the quantitative estimation of globulin may be effected. With this object the precipitate formed is treated in the manner described before when speaking of the estimation of albumin by weight (*Pohl*).²⁶⁵

[*Halliburton*²⁶⁶ adopts the following method:—The urine is neutralised and then saturated with magnesium-sulphate. If globulin be present, a precipitate forms. This precipitate may be collected on a filter and dissolved by the addition of water. The solution coagulates at 75° C. When the urine holds a large quantity of globulin, *Sir William*

Roberts' test will serve for its recognition :—A deep glass vessel is filled with water, and the urine is added, a drop at a time. Each drop leaves a milky track behind it ; and when much urine has been added, the fluid becomes opalescent, and clears again on the addition of acetic acid.]

5. Fibrinuria.—Fibrin is found in the urine in cases of hæmaturia and chyluria (*v. infra*). It then usually forms coagula. It is found, moreover, as a consequence of inflammatory exudation in the urinary passages. The coagula occur most commonly in croup and diphtheria, and occasionally in tuberculosis.

To detect fibrin the clots should be separated by filtration, washed with water, dissolved by boiling in solution (1 per cent.) of soda or (5 per cent.) hydrochloric acid (*Huppert*²⁶⁷), and the fluid tested, when cold, by the biuret test, or by Heller's test.

6. Hæmaturia.—The blood which occurs in the urine may be derived from several sources—from the kidney, renal pelvis, ureters, bladder, or urethra (*vide supra*).

In well-marked cases, the presence of blood is directly suggested by the colour of the urine, which varies from that of an extract of raw meat to a ruby-red ; but inasmuch as a similar appearance may be due to hæmoglobin in solution (hæmoglobinuria), it is never safe to base the diagnosis of this condition upon inspection alone. It may be accurately determined by—

1. *Spectroscopic Examination.*—The freshly-passed urine, which, if deeply coloured, should be diluted with water, will exhibit the two absorption-bands of oxyhæmoglobin (fig. 37), and these, on the addition of sulphide of ammonium, give place to the absorption-band of reduced hæmoglobin. Lastly, when blood containing urine has stood for some time, occasionally even in fresh urine, the spectrum of methæmoglobin may be looked for (fig. 42). •

2. *Heller's Test.*²⁶⁸—The urine is treated with caustic potash and boiled. The (basic) earthy phosphates are then precipitated, and together with them the hæmatin derived from the oxyhæmoglobin present. The phosphatic sediment is consequently coloured a bright red. Should it happen that the urine contains abundance of colouring matter (bile pigment, urobilin, &c.), which renders it difficult to appreciate the colour of the sediment, the latter should be separated on a filter and dissolved in acetic acid. The solution then becomes red if blood be present, and its colour vanishes gradually on exposure to the air.²⁶⁹

*Rosenthal*²⁷⁰ applies the test for hæmin directly to the dried precipitate (see Chapter I.). *Struve's*²⁷¹ method serves well for the detection of blood pigment. The urine is treated with ammonia or caustic potash, and the fluid then rendered acid with tannic and acetic acids. The presence of blood is shown by a dark-coloured precipitate, which, when

dried and treated with a little chloride of ammonium and glacial acetic acid, should yield *Teichmann's* crystals. This test is very sensitive, but less practicable than Heller's, which, with the spectroscopic examination, best meets the requirements of the physician.

3. *Almén's Blood Test.*²⁷²—A mixture in equal parts of tincture of guaiacum and mature oil of turpentine is poured on the surface of about 10 cc. of the urine. The presence of blood is shown by the appearance at the junction of the fluids of a ring at first white and afterwards turning blue.

If, in addition, the microscope reveals the presence of red blood-corpuscles, the existence of haematuria may be inferred, and it only remains to judge of its origin on the principles already laid down. For its clinical significance the reader may refer to this chapter: *Red Blood-Corpuscles*.

7. **Hæmoglobinuria.**—Sometimes the colouring matter of the blood is also found dissolved in the urine (see Chapter I.). This condition is apt to arise in the course of acute infectious diseases, in burns, and in various forms of poisoning. In the latter class of cases it is always a serious and even ominous symptom. It has been observed in poisoning by naphthol,²⁷³ carbolic acid,²⁷⁴ pyrogin,²⁷⁵ and chinin,²⁷⁶ and in erysipelas.²⁷⁷ Hæmoglobinuria is known as an idiopathic affection²⁷⁸ (paroxysmal hæmoglobinuria), and as such often occurs in connection with severe syphilis [and also in malaria and rheumatism, when the attacks appear to be determined by exposure to cold (*Taylor*).²⁷⁹]

The occurrence of hæmoglobinuria may be inferred when it is made evident by spectroscopic analysis and the application of Heller's and Almén's tests that the urine contains blood-colouring matter, whilst at the same time the microscope discloses smaller or larger masses of brown pigment, and either no red corpuscles, or so few as are inadequate to account for the results obtained. The spectroscopic appearances are usually those of methæmoglobin (fig. 42); and *Hoppe-Seyler*²⁸⁰ maintains that in this condition the blood pigment is always in the form of methæmoglobin.²⁸¹ [In seven cases of paroxysmal hæmoglobinuria investigated by *Halliburton*,²⁸² methæmoglobin was the only pigment present in three. In the remaining four there was found only methæmoglobin at first, but in a few hours the amount of pigment increased, and oxyhæmoglobin appeared as well. In such a specimen the spectra of methæmoglobin and of oxyhæmoglobin occur together. In three of Halliburton's cases serum-albumin was present in the beginning of the attack, and in one its appearance preceded that of the blood pigment. Albuminuria has been observed to alternate with hæmoglobinuria, and *Fagge* has described a paroxysmal albuminuria which he believed to be a mild form of the latter.]

8. Nucleo-Albuminuria.—The presence of small quantities of nucleo-albumin²⁸³ in the urine is not a pathological symptom ; when found in greatly increased proportion in women, it is often derived from the vagina. Such an increase, originating in the urinary passages, invariably points to catarrh of those parts. The urine in question is usually turbid when passed, and after a little while it deposits a bulky cloud. The latter is seen by the microscope to contain leucocytes and epithelium (*vide supra*). When a great quantity of mucin is passed, it forms a viscid gelatinous sediment at the bottom of the urine-glass, and no further evidence of its character is needed. *Fr. Müller*²⁸⁴ found large quantities in the urine of leukæmia, and *Obermayer*²⁸⁵ states that it is constantly present in jaundice. *A. Ott*²⁸⁶ has shown that a variable amount of nucleo-albumin occurs in every urine. According to *Mörner*,²⁸⁷ not nucleo-albumin but its antecedents are normal constituents of the fluid.

For the detection of nucleo-albumin chemically, the urine is treated with an excess of acetic acid, when it is rendered turbid if much nucleo-albumin is present. It may be necessary to dilute it previously, since concentrated urine, being rich in salts, will retain nucleo-albumin in solution even in presence of acetic acid. In testing for the presence of nucleo-albumin in albuminous urine, the great bulk of the albumin should be removed by boiling and filtering previously, and the filtrate allowed to cool before testing with acetic acid.

The best method for precipitating nucleo-albumin from urine is by the addition of acetate of lead (*vide supra*).

A. Ott's method for the detection of nucleo-albumin is very useful : to the urine is added an equal quantity of saturated salt solution, and *Almén's* tannin solution slowly supplied. If nucleo-albumin be present, even in very small proportion, an abundant precipitate will fall.

Almén's tannin solution is composed thus : tannin, 5 grms. ; 25 per cent. acetic acid, 10 cc. ; 40–50 per cent. ethyl alcohol, 240 cc.²⁸⁸

II. Carbohydrates.

1. Glycosuria.—Although various forms of sugar are occasional constituents of the urine, as, *e.g.*, sugar of milk after parturition, and lævulose (fructose) and maltose rarely, the consideration of these will nevertheless hardly detain us, since their manifestation, from the point of view of frequency and importance, possesses but little practical interest in comparison with that of the hexose²⁸⁹ grape-sugar (glucose, glycose, dextrose). Our remarks here will be confined to the occurrence and tests for the latter body.

(a) **Physiological Glycosuria.**—It must be mentioned at the outset that normal urine contains a trace of sugar. The fact of a physiological glycosuria was long ago laid down by *v. Brücke*,²⁹⁰ and has

quite recently derived remarkable confirmation.²⁹¹ *Wedenski* has made use of *Baumann's* discovery that benzoyl chloride forms insoluble compounds with the carbohydrates; and applying this to healthy urine, he succeeded in separating from the precipitate a body which gave all the reactions of grape-sugar. The proportion to be found in health, however, is so small, that it may be neglected as a disturbing factor, even in the most sensitive of the tests to be described. [This question of the presence of traces of sugar in the urine of health has been much discussed by English writers. According to *Sir G. Johnson*²⁹² normal urine is quite free from sugar; and *Mr. G. S. Johnson*²⁹³ has shown that when all the uric acid and kreatinin have been removed from such urine by precipitation with mercuric chloride, all reducing action disappears and no trace of sugar can be found. *Pavy*,²⁹⁴ by precipitation with oxide of lead, from large quantities of urine obtained the reactions of grape-sugar. His experiments, however, are the subject of controversy.²⁹⁵ *Halliburton*,²⁹⁶ in view of all the evidence at present available, inclines to the belief that normal urine does contain sugar; and *Allen*,²⁹⁷ using the same method as G. S. Johnson, also established its presence by the phenyl-hydrizin test.]

(b). Pathological Glycosuria.

(a.) Transitory Glycosuria.

1. *Spontaneous*.—Grape-sugar may appear temporarily in the urine in the course of many diseases, as cholera, intermittent fever,²⁹⁸ [typhus and typhoid], cerebro-spinal meningitis, and scarlatina,²⁹⁹ [after attacks of whooping-cough, asthma, and epilepsy (*Taylor*)³⁰⁰], in affections of the brain involving the fourth ventricle, in diseases of the heart, liver, and lungs, in gout, and in tertiary syphilis (*Ord*).³⁰¹ It has also been occasionally observed in small quantities in cirrhosis of the liver. Glycosuria in connection with these diseases is, however, very rare. It is commoner as an effect of certain poisons, notably morphia and carbon monoxide [after the inhalation of chloroform, ether, and amyl nitrite, and in poisoning with prussic acid, mercury, and curare. In some of these cases, however (morphia, curare, chloroform, &c.), it has been shown that the substance which reduces Fehling's solution in the urine is not sugar, but glycuronic acid (*Halliburton*)³⁰²]. The author has found grape-sugar in the urine in two cases of advanced asphyxia from breathing irrespirable gases (a mixture of carbonic acid and nitrogen).³⁰³

Feeding with thyroid gland causes carbohydrates to appear in the urine (*Eicard*,³⁰⁴ *Dening*³⁰⁵). In the author's hands, urine tested after the administration of the gland, by the ordinary sugar tests gave no result, while with Hoppe-Seyler's process it was often otherwise.³⁰⁶

2. *Alimentary or Induced*.—In certain diseases the assimilation of sugar is impeded (*Hofmeister*), and an alimentary glycosuria results.

This happens in hepatic cirrhosis (*Moritz, Kraus, Ludicig, Colasanti*) and in Graves' disease (*Chrostek*).³⁰⁷ Extensive disease of the brain of various kinds, but especially syphilitic, are attended by the presence of sugar in the urine (*Bloch, Strasser, v. Jaksch*).³⁰⁸ *Poll*³⁰⁹ found alimentary glucosuria in fever patients. *R. v. Jaksch*³¹⁰ has observed it in phosphorus poisoning, and only in those cases where the liver was profoundly affected; again in a case of yellow atrophy of the liver, and occasionally in connection with neuroses. Out of twenty-three cases of phosphorus poisoning that have come under the author's observation during the last few years, alimentary glucosuria was found in fifteen cases, that is to say, in 65.2 per cent. of the total. *v. Noorden*³¹¹ has shown that glycosuria is common in persons with fatty liver. Alimentary glycosuria arises in the course of pregnancy (*v. Jaksch, Lanz*).³¹² A consideration of these facts leads to conclusions which may sometimes be of service in diagnosis. Thus a transitory glycosuria arising in connection with traumatic neurosis may facilitate the recognition of the latter—in connection with phosphorus poisoning it points to serious involvement of the liver, and in an obscure case, to acute yellow atrophy. It may even take its place among the signs of pregnancy (*Lanz*),³¹³ and afford an early warning of impending diabetes (*v. Noorden*).³¹⁴

(β.) *Persistent Glycosuria*.—The continued excretion of grape-sugar in appreciable quantities belongs exclusively to diabetes mellitus, and it is the most certain symptom of that disease. Its great clinical importance lies in the fact that it ordinarily becomes apparent at a time when all other symptoms of diabetes are wanting. In such cases, however, one can be certain that diabetes exists only when, by repeated investigation, grape-sugar is found, and especially when its quantity is observed to increase with the administration of other carbohydrates, as cane-sugar,³¹⁵ and still better, starch.

Determination of Grape-Sugar.

(α.) *Qualitative Tests*.—It is very easy to determine the presence of a considerable proportion of sugar in the urine; but sometimes, when that body occurs only in traces or in very small quantity, the tests hitherto most commonly employed, namely, those of *Moore* and *Trommer*, are hardly sufficient for its detection. We have only recently become acquainted with a method which is in all cases adequate to the purpose.

1. *Moore-Heller Test*.³¹⁶—The urine is treated with liquor potassæ and boiled. If sugar be present, it is decomposed. Lactic acid and a number of volatile compounds are formed,³¹⁷ and with them certain coloured substances, which impart an intense deep-brown tint to the

fluid. This test is by no means accurate, and conclusions drawn from it are open to fallacy, since healthy urine turns brown with caustic potash from the action of that body upon nucleo-albumin. Moreover, the change of colour is proportional to the quantity of nucleo-albumin present, independently of sugar.

2. *Trommer's Test.*³¹⁸—The urine is rendered alkaline with caustic potash, and a fairly strong solution of cupric sulphate is added, drop by drop, until the cupric oxide formed ceases to be dissolved. The mixture is then heated in a test-tube. If sugar be present in greater quantity than a mere trace, a yellowish or red precipitate of the sub-oxide of copper falls before the boiling-point is reached, and at the same time the fluid loses colour somewhat.³¹⁹ This test is very sensitive. Trommer was able with it to detect sugar to the amount of 0.001, or even 0.0001 per cent. Unfortunately it is also ambiguous. The property of reducing cupric oxide in alkaline solutions belongs to a number of bodies which occur in healthy and morbid urine. Amongst these are uric acid, kreatin and kreatinin, allantoin, nucleo-albumin, milk-sugar, pyrocatechin, hydrochinon, and bile pigments. In addition to these, the ingestion of benzoic and salicylic acids, glycerine, and chloral leads to the formation in the system of substances which possess a similar reducing power. *Th. Lajon*³²⁰ observed that after the administration of sulphonal the urine reduces *Fehling's* solution. Hence it happens that the urine is sometimes thought to contain sugar on the evidence of Trommer's test when none can be found with other methods. The error is especially apt to arise when the boiling is continued for a long time. *The test can be depended on, therefore, only when reduction takes place at a temperature below boiling, which, however, occurs only when the urine contains a relatively large proportion of sugar.*

Fehling's fluid (*cide supra*) may be substituted for the copper sulphate and caustic potash in the process. [Pavy's³²¹ fluid, which is much used in England, is a modification of *Fehling's*. It has the following constitution:—(1) neutral potassic tartrate 640 grs., potassa fusa 1280 grs., water 10 oz.; (2) cupric sulphate 320 grs., water 10 oz. The two solutions are kept apart and mixed for use. Of the mixture 40 to 60 minimis are boiled in a test-tube, and a drop or two of the suspected urine is added. If heat be continued, the yellow precipitate shows itself in the upper part of the test-tube, and by adding more urine the fluid will be made to lose its blue colour entirely.]

A useful modification of Trommer's test has been suggested by *Worm-Müller*.³²² A mixture is made of 1.5 cc. of a 2.5 per cent. solution of cupric sulphate, and 2.5 cc. of an alkaline solution of tartrated soda and potash (prepared by dissolving 100 grms. of Rochelle salts in a normal solution of caustic soda), and heated to boiling-point; and in a

separate test-tube 5 cc. of the urine to be tested are boiled. The boiling fluids are added together without shaking them, and if sugar be present in any quantity, the suboxide is precipitated directly. Should no such precipitate form, the process is repeated with 2, 3, or 4 cc. of the cupric sulphate solution. This test is said to be very sensitive.

It should be mentioned that the power of a urine to dissolve cupric oxide does not necessarily imply the presence of sugar in the urine, since the property belongs to ammoniacal and albuminous urine, whether it contains sugar or not.³²³

3. *Fermentation Test.*—This depends upon the fact that grape-sugar decomposes in presence of yeast into alcohol, carbonic acid, and a number of other products (succinic acid, glycerine). It is conducted in the following manner:—A test-tube is filled for two-thirds of its depth with mercury, and in the remaining third with the urine, to which a little tartaric acid has been added. In this is placed some yeast which has been carefully washed. The mouth of the test-tube is then closed with the thumb, and it is inverted over a vessel containing mercury. If sugar be present, fermentation takes place directly, and the carbonic acid formed collects over the mercury. When yeast is placed in normal urine a limited fermentation takes place and some gas is disengaged. It is, therefore, well to compare the result of fermentation in saccharine urine with a healthy specimen (*Moritz*³²⁴). There are special fermentation tubes made for such purposes, and the experiment is greatly facilitated by their use.³²⁵ This test is sufficiently sensitive. It will serve to indicate 0.1 per cent. of sugar in the urine.³²⁶ [According to *Halliburton*,³²⁷ the fermentation test is the best for distinguishing sugar from other substances that reduce Fehling's solution in the urine. These substances are uric acid, hippuric acid, kreatinin, pyrocatechin, and glycuronic acid.]

4. *Phenyl-Hydrazin Test.*—In this we have a method for the detection of sugar which is greatly to be preferred to those already mentioned. Several years' experience has convinced the author that it is entirely accurate, simple in its application, and in every way suited to the needs of the physician. It depends upon the property of phenyl-hydrazin to form with grape-sugar a highly characteristic crystalline compound,³²⁸ called phenyl-glucosazone. This body has the form of yellow needles, and is but little soluble in water. The test is conducted in the following manner (*v. Jaksch*³²⁹):—Two parts of phenyl-hydrazin hydrochloride* and three of acetate of soda are placed together in a test-tube containing 6–8 cc. of urine. If the salts do not dissolve when the

* Twice as much of the phenyl-hydrazin salt as will lie on the point of the blade of a knife. [This salt is said to cause eczema, and should therefore be handled carefully.]

fluid is warmed, a little water is added, and the test-tube containing the mixture is placed for 20-30 minutes in boiling water. After this it is taken out and put into a vessel containing cold water. If sugar be present, even in very moderate quantity, there forms directly a yellow crystalline deposit, which may appear amorphous to the naked eye, but which when examined under the microscope is seen to contain yellow needles detached or arranged in clusters (fig. 134). If the urine holds but a very small proportion of sugar, the preparation should be placed in a conical glass, and the sediment examined carefully. In a case where only a mere trace of sugar exists, detached crystals of phenyl-glucosazone cannot fail to be seen. The discovery of smaller and larger yellow scales, or of powerfully refracting brown granules, must not, however, be mistaken for evidence of sugar. This test gives *very good*

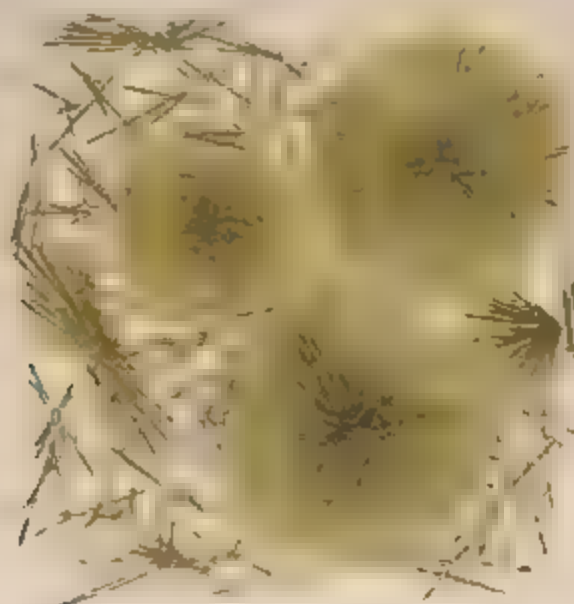


FIG. 134 Phenyl Glucosazone Crystals from Diabetic Urine
(eye-piece II ; objective 3A, Reichert).

results with every variety of morbid urine,³³⁰ and it is equally applicable whether albumin be present or not. In the former case, however, it is well to get rid of the excess of albumin previously by boiling. Phenyl-glucosazone crystals melt at 205°, and their character may be ascertained beyond doubt by submitting them to that temperature. The efficiency and utility of this test is amply proved.³³¹ By its means sugar to the extent of 0.1 per cent. can be detected. To verify the crystals their melting-point may be sought. Should this prove to be 205°, they may confidently be affirmed to be derived from a combination of phenyl-hydrazin with grape-sugar, and are conclusive evidence of the latter substance. The objections to its use made by *Geyer*, *Moritz*, and *Luther*³³² seem to be without weight. Doubtless the test requires much practice, but in expert hands highly satisfactory results are obtained.³³³

In addition to the foregoing methods, which especially merit attention, other tests for sugar have been suggested, and some of these call for description.³³⁴

5. *Böttger's Test.*³³⁵—A quantity of the urine is mixed with its own bulk of a concentrated solution of carbonate of soda, and a little basic nitrate of bismuth is added. The preparation is then boiled. If sugar be present, it turns black from the reduction of oxide of bismuth. This process has no advantage which does not belong to Trommer's test, and it is less sensitive. If the urine contains the principles of rhubarb taken as food, a black precipitate will fall independently of sugar.³³⁶ In albuminous urine, too, sulphide of bismuth will form a similar black deposit.³³⁷ These conditions must therefore be excluded.

The modification of *Böttger's* test by *Nylander*³³⁸ is more accurate. The reagent employed (called *Almén's* fluid) is prepared by dissolving 4 grms. of Rochelle salts (tartrated soda and potash) in 106 grms. of an 8 per cent. solution of caustic soda, warming the fluid, and adding as much basic pernitrate of bismuth as will remain in solution. The mixture so formed is added to the urine to be tested in the proportion of 1 in 11, and the whole is then heated. The fluid should blacken in the course of a few minutes. In this way it is asserted³³⁹ that sugar in the proportion of 0.1 per cent. can be detected. It has, moreover, the advantage of simplicity, but is also open to certain fallacies.³⁴⁰ It is inapplicable, as we have seen, to the case of albuminous urine, and the reaction occurs in presence of melanin or melanogen, or where the fluid contains a large proportion of reducing substances but no sugar.³⁴¹

6. *Rubner's Test.*³⁴²—The urine is treated with an excess of acetate of lead (sugar of lead), filtered, and to the filtrate ammonia is added until a permanent precipitate forms. The fluid is then heated, but not boiled. If sugar be present, the precipitate formed on the addition of ammonia gradually assumes a rose-red colour, which vanishes slowly on standing, more quickly on the application of heat (60°–70° C.), giving place to a yellowish coffee colour.

Rubner believes that the precipitate consists of sugar of lead. Milk-sugar does not give this reaction when the process is conducted as above, but when a solution of that body is boiled for 3–4 minutes with acetate of lead, and ammonia then added, a similar precipitate forms. In the author's experience, the best plan in performing the test is to heat the precipitate gradually at a temperature not exceeding 80° C. By its use *Penzoldt* has discovered sugar to the amount of 0.01–0.02 grm. in 10 cc. of urine. That writer employs a very simple and practical modification of Rubner's test.³⁴³ He adds to the urine a few drops of basic acetate of lead (subacetate) and a few of ammonia, and then warms the mixture.

The presence of sugar causes a red precipitate as before. This method is not less sensitive than the other.

7. *Mulder's Test.*—The urine is treated with carbonate of soda in solution, and solution of indigo-carmine is added until the whole is freely coloured. If sugar be present, the colour changes to yellow on heating, and becomes again blue when the fluid is shaken up with air.

This test may be conveniently applied thus.²⁴⁴—Two pieces of filter-paper are taken. One is placed in a concentrated solution of carbonate of soda, and the other in a solution of indigo-carmine. Both are then dried. When wanted for use, a small piece of the indigo carmine paper is placed in about 10 cc. of water, the urine under examination is added, and finally a large slip of the paper saturated with carbonate of soda is placed in the fluid. The result should be as above. The convenience of its application in this manner is the only merit of Mulder's test. It is neither sensitive nor accurate. [G. Ulmer²⁴⁵ recommends the use of indigo-carmine papers alone. The test solution is made from these with heat, and is again boiled with the urine. The colour obtained ranges from violet to straw-yellow, according to the quantity of sugar present. This is not a satisfactory test.]

8. *Johnson's Test (Picric Acid).*—Both Johnson and Thivry²⁴⁶ have employed picric acid as a test for sugar. A few drops of a solution of picric acid are added to the urine, which is then treated with caustic potash. If sugar be present, the fluid assumes a deep-red colour; but a red colour may also be obtained from caustic potash and picric acid alone, or in presence of kreatinin; and consequently, as a test for sugar, the process is not trustworthy.²⁴⁷ [Sir G. Johnson claims for picric acid the advantage that solutions do not change with keeping. When a drachm of normal urine is boiled with the same quantity of picric acid and half a drachm of caustic potash, a claret-red colour is produced, which in a test-tube of half-inch diameter will transmit light. This colour is due to kreatinin. The presence of the smallest quantity of glucose in addition will render the fluid so intensely dark that no light passes through.]

9. *Penzoldt's Test.*²⁴⁸—In this the reagent is diazobenzol-sulphonic acid dissolved in water, i. 60, without the aid of heat, but a drop of caustic potash may be added to facilitate solution. A few cubic centimetres of the urine under examination is placed in a test-glass, and rendered strongly alkaline with caustic potash. A like quantity of the reagent, which should be feebly alkaline, is now added. The process is repeated with healthy urine of the same tint and concentration, and the two specimens compared. Both are at first of a reddish-yellow colour, but whilst the healthy specimen remains unchanged, or nearly so, that containing sugar becomes a bright claret colour after some time. If sugar be present in abundance, the fluid eventually becomes dark red and opaque.

Penzoldt asserts that 0.1 per cent. sugar is appreciable in this way; but for practical purposes the test is not to be recommended, since acetone and diacetic acid have a similar reaction.²⁴⁹ Moreover, the substance used is highly explosive.²⁵⁰

10. *Möllisch's Reactions.*—Möllisch²⁵¹ has recently devised two methods, by the aid of which he believes that sugar can be detected in the urine, whether of health or disease.

(a.) The first depends upon the reaction of sugar with α -naphthol and sulphuric acid. To obtain this, he takes $\frac{1}{2}$ to 1 cc. of the fluid containing sugar—urine should be highly diluted for the purpose—places it in a test-tube, and adds to it two drops of a 15-20 per cent. alcoholic solution of α -naphthol. The fluid becomes turbid from the precipitation of some of the α -naphthol. Concentrated sulphuric acid is now added in excess, and the whole is well mixed.

The presence of sugar is shown by the transitory appearance of a blue colour, and the formation of a violet-blue precipitate on the subsequent addition of water.

(b.) This reaction is obtained with thymol and sulphuric acid. The urine which is thought to contain sugar is highly diluted, and to $\frac{1}{2}$ 1 cc. in a test tube is added first two drops of a 15-20 per cent. alcoholic solution of thymol, and the sulphuric acid in excess. When the mixture is shaken, the momentary development of a "cinnabar-red-carmine red" discloses the presence of sugar, this colour giving place to carmine when the fluid is diluted with water.

By this method Molisch maintains that so small a proportion as 0.00001 per cent. of sugar can be detected. Similar reactions, however, may be obtained with cane-sugar, fruit-sugar, and maltose. Seeger³⁵ has further investigated the subject, and found that chemically pure solutions of proteids, and especially of serum albumin, behave in like manner. The author has repeatedly performed the test with albuminous urine, with the result that the α -naphthol reaction was obtained when the fluid was diluted in a much higher degree than that (1:100) recommended by Molisch in the case of sugar. In presence of albumin, the dark-violet coloration of the fluid was followed by the deposition of a greenish-black precipitate. The thymol and sulphuric acid reaction with this body was almost identical with that displayed by sugar.

For these reasons, Molisch's reactions, which no doubt find a valuable application in vegetable physiology, are not to be recommended as a test for glycosuria. The researches of Mylius and c. Udransky³⁶ have made it clear that Molisch's reaction is identical with the furfural reaction, which takes place not only with sugar, but with any carbohydrate.

11. *Hoppe-Seyler's Test.*³⁷ To 5 cc. of a 0.5 per cent. solution of orthonitrophenyl-propionic acid in caustic soda and water, are added ten drops of urine, and the mixture is boiled for fifteen seconds. If sugar be present, a blue colour develops (formation of indigo). The reaction is not impeded by the presence of albumin, but it is yielded equally by levulose, maltose, and sugar of milk. The test has the advantage of being applicable to a small quantity of urine, and is useful provisionally, but is not entirely to be depended on. If, when the fluid is cool after boiling, chloroform be added to it, the latter will take up the indigo derived from sugar that may be present in the urine and assume a beautiful blue tint.

The author finds, from an extensive series of experiments, that this test does not afford a reliable means for the detection of grape-sugar, since urines, proved by reliable tests (Trommer's test; fermentation; phenyl-hydrizin) to be free from sugar, will give positive results with Hoppe-Seyler's reagent.

12. *Resorcin Test.* — E. Fischer and W. L. Jennings³⁸ have used resorcin as a test for the carbohydrates (grape-sugar, maltose, cellulose, &c.) in small quantities. If much sugar be present, the urine should be diluted. To 2 cc. of urine 0.2 gm. resorcin is added in a test-tube, which is put on ice to cool, and hydrochloric acid gas is led into it. The fluid is allowed to stand for several (twelve) hours at the ordinary temperature, is then diluted with water saturated with caustic soda, and treated with Fehling's solution and warmed. A reddish-violet colour attests the presence of carbohydrates. Normal urine is never free from these substances, and it therefore yields the reaction.

It should be remarked, in addition, that Fröhlich's³⁹ methylene-blue method, which was tested by Hoke in the author's clinic, has not proved reliable for the detection of grape-sugar in urine. Bremner⁴⁰ recommends the employment of aniline dye-stuffs for detecting sugar in the blood and in urine, the diagnostic value of this certainly interesting report still requires, however, to be tested on a sufficiency of material.

It remains to mention that experiments have been instituted by several investigators (*v. Brücke, Seegen, Abeles, Salkowski*³⁵⁸) with the object of separating the sugar when it exists in very small quantities in the urine, and submitting it in the concentrated form to the action of Trommer's, the phenyl-hydrazin, and other tests.

The separation and detection of grape-sugar, as of the carbohydrates generally, may be effected by the action of benzoyl chloride, which with the carbohydrates forms insoluble benzoic acid æthers. For this purpose the urine is treated with benzoyl chloride and caustic potash. If to a litre of urine there be added 200 cc. of a 10 per cent. soda solution and 10 cc. of benzoyl chloride, the nauseous smell of the latter disappears on shaking the mixture, and a precipitate results. The addition of concentrated sulphuric acid and a few drops of an alcoholic solution of α -naphthol causes an intense red colour if a trace of benzoyl combination with the carbohydrate be present, and the coloured fluid will exhibit a well-defined absorption-band in the green part of the spectrum.³⁵⁹

It is indispensable that both the sulphuric acid and the naphthol solution should be absolutely pure. To ascertain that they are so, a 10 per cent. solution of α -naphthol in chloroform is prepared, and to a drop of this in a test-glass is added first 0.5 cc. of water, and then 1 cc. of pure sulphuric acid. If the reagents are serviceable, the mixture assumes a yellow tinge. A little is next added to the fluid to be investigated—*i.e.*, to a benzoic acid æther precipitate suspended in water. A reddish-violet ring is evidence of sugar or a carbohydrate (*Luther, Roos*).³⁶⁰ This proceeding, though simple, is open to fallacy, since many other substances, as albumin, fats, &c., form acid æthers with benzoyl chloride. In other words, it is a form of the furfural reaction, and partakes of its ambiguous character. The necessity of securing absolutely pure reagents is a further disadvantage which detracts from its value as a clinical test.

(β.) Quantitative Estimation of Grape-Sugar.

1. *Titration Method (Fehling)*.³⁶¹—The principle upon which this method is founded depends upon the property which grape-sugar possesses of reducing cupric oxide to its suboxide in alkaline solutions. It has been applied in various ways, and most of these are to be found described in the text-books of urinary chemistry.

Accurate results are obtained by the titration method in the following modification proposed by *Soxhlet*.³⁶²

(A.) Preparation of the Solutions.

In the first place, the Fehling's solution—which must be *freshly* prepared before each titration—is compounded from the following constituents:—(1) A solution of copper sulphate; (2) a solution of Seignette

salt (potassium-sodium tartrate); (3) a solution of caustic soda. The Fehling's solution should contain 34.64 grms. of crystallised copper sulphate per litre.

1. *Preparation of the Copper Solution.*—A saturated solution of copper sulphate is prepared by adding fresh quantities of the chemically pure salt to one and the same portion of hot water so long as any continues to be dissolved. The solution is then filtered whilst still hot, and the filtrate is continually stirred as it cools, a blue crystalline pulp of copper sulphate being deposited during this operation. The crystals after being drained on a filter are spread out on a wide plate and set aside in a dry place. When sufficiently dried—a condition indicated by their whitish-blue colour—the crystals are packed in a dry glass vessel and stored for use.

To ensure accuracy it is advisable to determine the water content of the crystals by drying a sample at $100^{\circ}-110^{\circ}$ C. If of the proper degree of dryness, 100 grms. of the crystals will lose in weight during this operation to the extent of 28.87 per cent.

A quantity of these crystals, weighed in a chemical balance, is dissolved in a little hot water, the solution being transferred to a measuring glass, the beaker rinsed out carefully with water, and the contents of the glass diluted until the solution contains 103.92 grms. of copper sulphate per litre. It is then stored for use in a bottle closed by a caoutchouc stopper.

2. *Preparation of the Seignette Salt Solution.*—The pure commercial salt (potassium-sodium tartrate) is weighed out in an ordinary balance and dissolved in hot water, the solution being filtered and afterwards diluted with sufficient water to reduce the content to exactly 280 grms. of tartrate per litre.

If, for example, 140 grms. of Seignette salt were dissolved, the solution must be made up to 500 cc.

Since this solution is liable to become infested with mould, it is advisable to add a little carbolic acid.

3. *Caustic Soda (Sodium-Hydrate) Solution.*—This solution (sp. gr. 1.137) must be clear, the clarification being preferably effected either by standing or by filtration through asbestos.

When this titration method is in constant use it is well to have a stock of 1 litre of each of the above solutions ready at hand.

(B.) *Performance of the Method.*

The Fehling's solution for the day's consumption is prepared, immediately before commencing operations, by *accurately* measuring out, in order, into a measuring vessel fitted with a ground stopper, equal volumes of the solutions described above, e.g., 50 cc. of No. 1, 50 cc. of No. 2,

and 50 cc. of No. 3. When shaken up, the mixture forms a perfectly clear Fehling's solution, 1 cc. of which corresponds to 0.005 grm. of grape-sugar.

150 cc. of the above solution contains 5.196 grms. of copper sulphate, consequently 1000 cc. will contain 34.64 grms., the exact amount of copper sulphate required in Fehling's standard solution.

The approximate sugar content of the urine under examination being ascertained by means of a specific gravity determination, 10 cc. of the Fehling's solution are measured out from a burette and mixed with 0.5 cc. of urine from a second burette, together with 40 cc. of water and a few drops of strong caustic soda, the mixture being then heated. Should the liquid still retain its blue colour, a further quantity of urine is added, but if it is already decolorised a fresh test is made with a smaller quantity of urine—0.4 cc., &c.

Assuming the result to show that 10 cc. of Fehling's solution are exactly decolorised by 1 cc. of the urine under examination, then, since this quantity of the reagent corresponds to 0.05 grm. of grape-sugar, 1 cc. of the urine contains 0.05 grm. and 100 cc. contain 5 grms. of grape-sugar. Performed, however, in the above manner the determination is not perfectly accurate, and if precision is desired, 5-10 cc. of the urine must be employed to reduce 10 cc. of the Fehling's solution, the urine being diluted, in accordance with the sugar-content indicated by the preliminary experiment, so as to contain from 0.5 to 1 per cent. of sugar. Thus, in the example previously cited, the urine would require to be diluted to a tenfold volume. With this object the urine is placed in a burette, and 10 cc. are run into a 100 cc. measuring glass, which is then filled with water exactly up to the 100 cc. mark. Since in the preliminary experiment 1 cc. of the undiluted urine was required to effect the reduction of 10 cc. of Fehling's solution, the final test—assuming the preliminary experiment to have been correctly performed—may require from 9.5 to 10.5 cc., consequently ten 100 cc. flasks are charged with 10 cc. of Fehling's solution, 40 cc. of water, and a few drops of caustic soda in each, 9.5 cc. of the dilute urine being then added to the first flask, 9.6 to the second, and so on through the series. The ten flasks are then placed together on a sand-bath and heated to boiling, after which they are simultaneously removed and placed on a white support.

A sand-bath of suitable dimensions which the author has had made specially for this purpose answers admirably, enabling more than twenty flasks to be heated at a time.

Suppose, for example, the liquid in the flask containing 9.8 cc. of urine still remains blue, whilst that with 9.9 cc. has become decolorised;

then in this case 9.9 cc. of the tenfold diluted urine has been consumed in reducing 10 cc. of Fehling's solution. Now, these 10 cc. of Fehling's solution correspond to 0.05 grm. of sugar, consequently 9.9 cc. of the diluted urine, or 0.99 cc. of the original urine, contain 0.05 grm. of sugar, and therefore 5.05 grms. of grape-sugar are present in 100 cc. of the urine.

When performed in the foregoing manner, devised by Soxhlet, the method yields excellent results, agreeing completely—according to the author's extensive personal experience—with those obtained by the aid of a good polarimeter. All other modifications of the titration method, as also that proposed by *Leube* and *Salkowski*,³⁶³ are far less accurate. The Allihn-Soxhlet method, which was devised for solutions of pure sugar, is inapplicable to urine, since the ammonia always present in urines dissolves copper suboxide, and, consequently, renders an exact determination impossible; hence the observations that have been made by the aid of this method must be set down as inexact.

When one is well skilled in the performance of the Soxhlet method, it is very rapid, so that in a brief space of time a very large number of titrations can be made, all of which may be considered as perfectly accurate. Furthermore, the method can be advantageously utilised for the quantitative determination of the pentose group of sugars, to be dealt with on a subsequent page (see p. 337). In cases where the urine contains glucose in addition to one or more members of this group, the concurrent determinations made by the polarimeter and by titration will give the values for both sugars.

[*Mr. Martindale* recommends the following formula:—(1.) Sulphate of copper, 181 grains; distilled water to 6 ounces. Dissolve. (2.) Tartrate of potassium, neutral, 728 grains; caustic soda, 360 grains, distilled water to 6 ounces. Dissolve.

Of a mixture of these two solutions (Fehling's fluid) in equal volumes, 10 cc. will be decolorised and reduced by 0.05 grm. of glucose or diabetic sugar in solution.

Titration with Pary's Ammoniated Cupric Solution—This is a fluid composed as follows:—Cupric sulphate, 4.158 grms.; potassic sodic tartrate, 20.4 grms., caustic potash, 20.4 grms., strong ammonia (sp. gr. 0.880), 300 cc.; water, 1 litre. The object of the ammonia is to prevent the precipitation of the suboxide, and so to render the decoloration of the copper solution more evident. In preparing the fluid, the caustic potash and the tartrate are dissolved together, the cupric sulphate by itself; the two solutions are mixed, and when cold the ammonia is added, and water supplied to the specified bulk. Ten cc. of this solution are decolorised by 0.005 grm. of sugar. *Pary* recommends for clinical convenience the use of hermetically sealed tubes of glass, each containing 10 cc. of the solution.^{364]}

2. *By Fermentation.*—This method of analysis was first suggested by *Roberts*,³⁶⁵ and *Worm-Müller*³⁶⁶ applied it to determine the proportion of sugar in the urine from the density of that fluid before and after fermentation. According to *Worm-Müller*, this method with the aid of a thermometer and a pycnometer fitted with a graduated index-scale, gives good results where so little as 0.5–1 per cent. of sugar is present. *Roberts* concluded from experiments that a difference of 0.001 sp. gr. corresponded to 0.23 per cent. of sugar, and arrived at the following formula :—

$$x = \frac{D \times 0.230}{0.001}$$

Where

• x = the percentage of sugar.

D = the difference in the density of the urine before and after fermentation.

It is possible to obtain good approximate results from the application of this method clinically. The following apparatus is required: Two hydrometers accurately graduated to four places of decimals, and each provided with a thermometer carrying a fractional index, and capable of registering $\frac{1}{10}^{\circ}$ C. These instruments should measure densities ranging between 1.000–1.025 and 1.025–1.050 respectively up to four places of decimals.

The first step is to take the sp. gr. of the urine at the temperature for which the hydrometer in use was constructed. This may be done by placing the test-glass containing it in a vessel of water which may be cooled or heated as required. 100–200 cc. of the urine are then placed in a flask, together with some fresh yeast which has been carefully washed on an ash-free filter to secure removal of inorganic impurities. The flask is next closed with the arrangement represented in fig. 135, by which evaporation is prevented. Fermentation is allowed to go on for 24–48 hours. After this, the fluid, which should be clear, or nearly so, is passed rapidly through a filter, and its density again taken at the appropriate temperature, the latter being secured and ascertained as before. From the observations made, the percentage of sugar is calculated by the formula given above.

In twelve cases of diabetes this method proved most valuable. Its simplicity renders it very suitable for clinical use.

*Th. Lohnstein*³⁶⁷ has lately introduced an aerometer which makes the test simpler of application; clinical experience of this is as yet wanting.

The figures in the following table afford the means of comparing the results obtained by fermentation and by means of the polarimeter in a series of cases :—

	Percent.						
1. Fermentation:	2.22	3.55	4.49	5.38	6.06	6.23	6.0
2. Polarimetry:	2.25	3.65	4.67	5.60	6.01	6.00	6.1

³⁶⁸

In the polarimetric investigations the instruments of *Lippich* and *Venzke-Soleil* were used. It appears from the table that in seven instances approximately identical figures were obtained. Nothing more is needed to enforce the value of the fermentation process as a clinical test.

3. *By Polarisation.*—Grape-sugar possesses a dextro-rotatory power for polarised light, and upon this fact a quantitative test for that body is based. Such a test must necessarily be open to fallacy, inasmuch as the urine in diabetes is apt to contain other substances, such as β -oxybutyric acid and levulose, which rotate light in the opposite direction. It is well, therefore, in applying it, to do so both before and after fermentation (*Hoppe-Seyler*, *Kutz*, *Worm Muller*, and *K. A. H. Mörner*), when the difference in the results will be a measure of the quantity of grape-sugar in solution. The advantage of the method is that it dis-



FIG. 135.—Flask for the Approximate Estimation of Sugar by Fermentation (4).

penses with delay. In recent times it has been invested with great accuracy by the use of a polarimeter constructed on the principle devised by *Lippich*. This instrument is represented in fig. 136.³⁰⁹ It is employed thus:—

The graduated disc is turned towards the observer, and the tube towards the lamp, and the cap which covers the telescope and the posterior end of the apparatus is removed. The lamp is placed at a distance (45 cm.) equal to the length of the instrument itself. Carbonate of soda is placed in the cage attached to the lamp, and fused until the receptacle is full. The latter is then fixed in the lamp in such a way as to be in contact with the flame at the side only, and a screen is so placed that the light passes only through the aperture in it.

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FIG. 30. Lippich's Polarimeter ($\frac{1}{2}$).
 a. Astronomical telescope.
 b. Analyser (Nicol).
 c. Half prism (Nicol) fixed (polariser).
 d. Complete prism (Nicol) movable (polariser).
 e. Illuminating lens.
 f. Diaphragm.
 The diagram shows a horizontal section of the instrument as seen from above.

The following description will sufficiently explain the construction of the instrument:—At the hinder end of the polarimeter is a rod supporting the arc of a circle, and behind this, in the direction of the lamp, another rod, with a notch marked upon it. The second rod is fitted with a screw, by means of which it moves upon the first, and can be fixed in contact with the latter. When the line upon the upper surface of the adjustable rod corresponds to the central point (*o*) of the metallic arc, and this with the *o* of the vernier, the entire field is dark, and by rotating the arc (which is done with the ivory lever in front) both halves will be obscured or illuminated equally. In experimenting with the instrument, the adjustable rod is deflected slightly to the right or to the left. It is practically a lever attached to the box containing a Nicol prism (*d* in the section, which is represented as seen from above), and rotates the prism on its axis.

In conducting an observation, the tube containing the urine or other fluid to be examined is placed in the capsule, and the second rod is moved right or left, say as far as the mark 4. The observer then looks through the telescope, and adjusts the instrument in such a manner that the field is illuminated to the utmost in one-half at least; after which he moves the telescopic tube until the vertical line dividing the field into two equal parts is rendered as narrow and well defined as possible, and the position of the flame is again looked to. The ivory lever is now moved forward, and the *inner notched* rim of the disc is rotated to the right or the left, until both halves of the field are equally obscure. The lever is next reversed, and the observer works the micrometer screw at the lower part of the disc, whilst he looks for a difference in the degree of illumination of the two halves of the field. If this does not occur, the latter was either too bright or too obscure. It may be rendered brighter by increasing and inverting the angle between the two rods (*c*, *d*) to which the polarisers are connected; and the less this angle, the less will be the variation with equal differences of adjustment. When the proper degree of illumination is obtained, a number of readings are taken with different adjustments.

The number of degrees to the nearest $\frac{1}{2}^{\circ}$, from the zero-point of the disc to that of the vernier, are read off; and then, proceeding in the same direction, the first line on the vernier which exactly corresponds with a division on the disc is noted. This is easily found by inspecting the divisions to the right and left of that one which is assumed to be correct. Both are on the *inner* side of the corresponding marks on the circle. The longer lines on the vernier correspond to 0.01° , the short ones to 0.005° . In practice, the adjustments should not differ by more than 0.005° .

The way in which the readings are taken may best be shown by an

example :—Supposing the zero-point of the disc stood to the right of the zero-point of the vernier, and that, between the two, $\frac{3}{4}^{\circ}$ in addition to twenty long vernier marks and one short one had been counted, then $+ \frac{3}{4}^{\circ}$, 205 is written down. With further readings, the vernier marks only are noted, added together, and the average taken. Should the result again be 205, then

$$\frac{3}{4}^{\circ} = 0.75^{\circ}, 0.75^{\circ} + 0.205^{\circ} = + 0.955^{\circ}.$$

One long mark = 0.01° , 20 long marks = 0.200° .

One short mark = 0.005° , 1 short mark = 0.005° .

Finally, the zero-point is fixed by taking the tube out of the capsule *without otherwise altering the arrangement of the instrument*.

The field of vision is unequally illuminated, and the mark no longer distinct. The telescope is focused to show the mark, the lever brought forward, the disc put in with the hand, the lever reversed, and the micrometer screw carefully adjusted. A number of readings is taken, and an average struck. Supposing this to be $- 2.045^{\circ}$ (*i.e.*, the zero-mark for the present inclination of the polariser rods), the number is to be deducted from the original reading; thus, $+ 0.955 - (- 2.045) = 0.955 + 2.045 = 3.0^{\circ}$.

If the urine was examined in a tube two decimetres long, then

$$2 [\alpha]_D = 3.0^{\circ}, \text{ and } [\alpha]_D = 1.5.$$

$[\alpha]_D$ being the specific rotating power for grape-sugar ($C_6H_{12}O_6$), is in this case = $+ 52.5^{\circ}$.

52.5° for 100 grms. in 100 cc.

: 1° for $\frac{100}{52.5}$ grms. in 100 cc.

1.5° for $\frac{100 \times 1.5}{52.5}$ grms. in 100 cc.

The percentage of sugar in the case taken would therefore be

$$\frac{100 \times 1.5}{52.5} = 2.85 \text{ per cent.}$$

Care should be taken throughout the observation that the position of the lamp is not altered, otherwise varying results are obtained. During the whole process, therefore, the same position of the lamp and instrument must be maintained, and all readings should be taken continuously with one filling of the platinum cage.

By carefully observing these precautions, very accurate results may be obtained.

The instrument will serve also to show whether a liquid has the power of rotating light or not. In the latter case, the zero-point of the disc stands in the same lateral position to the zero-point of the vernier as the second pointer to the zero-point of the segment connected with the polariser (right or left), the deviation of the zero-point of the disc from that of the vernier being half as great as the deviation of the two rods.

2. Fructosuria.—Fruit-sugar (*lævulose*) is found sometimes in conjunction with grape-sugar in the urine (*K. Zimmer, Seegen, May*).³⁷⁰ The latter will then yield all the reactions of grape-sugar, including that with phenyl-hydrazin. It may happen under such circumstances that a specimen of the urine examined with the polarimeter will fail to rotate polarised light to the right, or may even rotate it to the left; and from this fact the presence of *lævulose* may be inferred.

Lævulose taken as food by diabetics is partly absorbed by the system (*Külz, Haycraft, Palma*),³⁷¹ and, according to *Haycraft* and *Palma*, it is eliminated in part as such and partly as glucose. When administered to diabetics it does not appear in the urine. Observations by *r. Stransky*,³⁷² in the author's clinic, serve to show that it is changed into other carbohydrates, and especially into grape-sugar. This appears to occur also in certain cases of abdominal tumour.

3. Lactosuria.—Milk-sugar occurs in the urine of women who are nursing.³⁷³ Its recognition can be effected only by separation from the fluid (*Hofmeister*). An effort has been made with the phenyl-hydrazin method to transform this body in the urine into phenyl-lactosazone (*r. Jakob*,³⁷⁴ but as a test the expedient has not succeeded.

The presence of milk-sugar may be inferred if, while no result is obtained with the phenyl-hydrazin and the fermentation tests, evidence of sugar is given by *Trommer's* and *Nylander's* tests, but only after prolonged boiling. *Rubner's*³⁷⁵ test for distinguishing grape-sugar from other carbohydrates is as follows:—The urine is treated with solid acetate of lead, well boiled, and ammonia added, when a rose colour develops. The colour in the case of grape-sugar is a coffee-brown; with chemically pure maltose light yellow; while *lævulose* displays little or no change.

In a case of traumatic neurosis where the administration of grape-sugar was followed by the appearance of that substance in the urine, milk-sugar had a similar result. The test just described gave a reaction, and the urine yielded right-sided polarisation. Experiments made with alimentary lactosuria have shown that the process is a good one for the detection of lactose. To distinguish lactose from glucose, *Ruizard*³⁷⁶ proceeds to decompose by hydrochloric acid, pass the constituents into osazone (see this chapter: *Phenyl-hydrazin Test*), and filter while hot. Glucosazone will remain on the filter; while galactosazone crystallises out in the cold, and can be recognised by determination of the melting-point (188°–191° C.).

The same observer advocates another method. Acetate of copper is reduced by grape-sugar, and not by milk-sugar. A solution after treatment with hydro-

chloric acid (formation of glucose and galactose) will be reduced on the application of heat. To apply this test to the urine, the latter is heated with hydrochloric acid and then tested as above. In a recent case within the author's personal experience, however, both these methods proved unreliable.

4. Dextrin. Dextrin has been seen in the urine of diabetics (*E. Reichard*³⁷⁷), where it seemed to take the place of grape-sugar. In such cases Reichard found that the urine behaved with Trommer's test in all respect like a solution of dextrin, the originally blue fluid becoming first gradually green, then yellow, and sometimes dark brown.

5. Animal Gum.—*Landwehr*³⁷⁸ has recently found in the urine a carbohydrate presenting a close analogy to members of the gum series. To this he has given the name of "animal gum," and he believes that it is a normal constituent of the fluid. The methods for its detection and isolation will be found in the original contribution on the subject. The statements there made are confirmed by *Wedenski*³⁷⁹. Observations by *Haycraft* and *Palmer*³⁸⁰ showed that maltose when administered to diabetics is eliminated partly as glucose. The presence in the urine of this substance is not evidence of pancreatic disease. In affections (as, for instance, carcinoma) of the pancreas the administration of carbohydrates is not always followed by their appearance in the urine. The administration of glucose is not followed by maltosuria, and this condition is wanting in typical pancreatic disease. On the other hand, the author has observed maltosuria occasionally in various disorders, including malignant tumours. It may be mentioned that sometimes cane-sugar taken as food reappears as such in the urine.

Amongst other carbohydrates occasionally present in health and disease are maltose (*Le Nobel*, *r. Ackern*³⁸¹). A left-rotatory carbohydrate was observed by *Leo* and *Kulz*³⁸² in diabetic urine.

6. Pentose—This substance, in the form of arabinose, rhamnose, and xylose, has been identified in the urine by *Salkowski*³⁸³. According to the author's personal observation, it appears that pentoses are frequently contained in beer. Probably, therefore, beer is the source from whence the small quantity of such sugars he has not infrequently found in urine is derived.

When the urine contains **Arabinose**, it will deflect the plane of polarised light towards the right, and will also give positive results with Trommer's and Nylander's tests, as well as exhibiting typical crystals with phenyl-hydrizin, but is *not* fermented by yeast. Such urine also answers to the Tollen's interruption test,³⁸⁴ i.e., when boiled with hydrochloric acid of sp. gr. 1.19 the urine assumes an intense cherry-red coloration, and deposits on cooling a dark flocculent precipitate, which, when filtered off, can be dissolved to a violet solution in alcohol. This solution when examined in a spectroscope exhibits a

very characteristic band between Frauenhofer's lines D and E of the spectrum.

Xylose behaves in the same manner, except that the dextro-rotation is much less extensive.

Rhamnose does not answer Tollen's interruption test, and furthermore gives a small dextro-rotation of the plane of polarised light.³⁸⁵

They have been found in connection with the morphia habit (*Salkowski, Justowicz, Reale*³⁸⁶), in diabetes (*Katz and Vogel*³⁸⁷), and in apparently healthy persons (*Blumenthal*³⁸⁷). *Lindemann and May*³⁸⁸ have ascertained that when rhamnose is administered about 8 per cent. is eliminated in health and about 16 per cent. in diabetes. In the course of researches conducted by the author it was found that in one case of diabetes 37.08 per cent. of the arabinose administered to the patient was excreted in the urine and 11.9 per cent. in the faeces, 51.02 per cent. thus having been absorbed into the system. In non-diabetic cases from 52.7 to 98.8 per cent. of the arabinose administered was not recovered. Xylose was absorbed to the extent of 42.8-73.3 per cent. by non-diabetics. In the case of rhamnose, 68.80 per cent. disappeared in non-diabetics, and 74.62 per cent. in one diabetic case, 82.09 per cent. being eliminated in a second instance (*R. v. Jaksch*³⁸⁹).

III. Choluria. Of the bile constituents, the biliary acids and pigments chiefly concern us here. A third constituent of the bile, cholesterol, has never yet been found in the urine in jaundice, although it occurs in considerable quantities in other connections.

*Hoppe-Seyler*³⁹⁰ has satisfactorily proved that biliary acids occur in the urine of jaundice; but their presence is of relatively little clinical interest, since they can be detected only by tedious chemical processes, but seldom suited to our purpose. None of the methods which have been suggested for the recognition of these bodies directly in the urine can be relied upon. That of *Mackay*, which is founded upon their physiological properties, is perhaps the most useful with which we are acquainted. Where the presence of bile acids in considerable proportion is suspected, resort may be had to the method for their detection in the blood (p. 91). The biliary acids, when isolated, or obtained as an alcoholic extract from the evaporation residue of the urine, may also be submitted to the furfural test. To this end the fluid is treated with a few drops of a 0.1 per cent. watery solution of furfural and sulphuric acid. The presence of biliary acids is shown by a red coloration³⁹¹. This reaction, however, is given by so many substances that its value as a test is slight (see *supra*).

A fact of much greater practical interest is the occurrence of bile pigments in the urine. This results from the comparative ease with which they may be recognised. Their manifestation implies, in the

first place, that there has been obstruction of the bile-ducts in the liver, in consequence of which substances secreted by that organ have made their way into the lymphatics and the general circulation, from which they are subsequently eliminated by the kidneys. This is *hepatogenic jaundice*, the commonest form of choluria. The conditions which cause it are manifold and various. Of these, simple obstruction or narrowing of the bile-ducts is the most obvious. But when it is remembered that the pressure under which the bile is secreted is very slight, it will become apparent that other circumstances—as, for instance, one-sided immobility of the diaphragm, thrombosis of the portal vein, &c.—will tend to retard the propulsion of bile, and consequently to induce choluria. Such causes must be discriminated by physical examination, and on other grounds.

The bile pigments of urine are not in all cases necessarily derived from the liver. It is conceivable that, with a quite normal biliary function, these bodies may be present, and they then owe their origin to blood-colouring matter which has undergone certain changes (see Chapter I.); and these changes may take place either in the blood proper (*haematogenic jaundice*³⁹²), or supervene when the colouring matter has been discharged in the tissues (*Quincke's inorganic jaundice*).³⁹³

It follows from what has been said, that the appearance of bile pigment in the urine is a fact of very extended import, and caution must be observed in inferring from it the existence of hepatic disease. Still the latter is by far its more common association.

Urine containing bile pigments is usually clear, deeply stained a yellow- or greenish-brown, and when shaken shows a yellow froth even when these bodies are in very small proportion. The chemical tests for bile pigments are very numerous, but it will suffice here to describe three, which alone, perhaps, are entirely trustworthy.

It has been claimed for the “cholecyanin test” of Stokvis³⁹⁴ that it is the most sensitive of all; but of this the author has no experience. It must be borne in mind that bilirubin alone, of all the bile pigments, occurs in fresh urine; the other pigments, biliverdin, bilifuscin, and biliprasin, being its oxidation products.

1. *Gmelin's Test.*³⁹⁵—A small quantity of nitric acid containing nitrous acid is placed in a test-glass, and a little of the urine under examination is poured in a separate layer upon its surface. To do this, care will be necessary, and it will be facilitated by slanting the test-glass so that the urine shall float on the surface of the acid. If bile pigment be present, there is a play of colours at the point of contact between the acid and the urine, and a green ring in particular forms, indicating the production of biliverdin. Gmelin's test will not serve for urine which has been treated with alcohol, since that body, when

similarly brought in contact with nitric acid, also yields a beautiful bluish-green ring (*H. Huppert*).³⁹⁶

In *Rosenbach's*³⁹⁷ modification of Gmelin's test, the urine is passed through a filter, and on the latter a drop of nitric acid is allowed to fall, when the coloured rings will develop around it. This process affords a very sensitive test, but its results can be relied upon only when the filter-paper employed is known to be absolutely free from impurities (colouring matter) which might account for them.

*Dragendorff's*³⁹⁸ method is to place a little of the urine on a plaster of Paris disc, and when the greater part has been absorbed, a drop of nitric acid is applied to the remainder. A parti-coloured ring forms around, in which the green tinge is prominent.

2. *Ultzmann's test*³⁹⁹ is very serviceable in cases where the urine contains a considerable proportion of bile pigment. The urine is mixed with solution of caustic potash (one part to three of water) in a test-glass, and hydrochloric acid added. The production of biliverdin by oxidation is indicated by the fluid assuming an emerald-green tint.

3. *Huppert's method*⁴⁰⁰ serves for the detection of the merest traces of bile pigments. 8 to 10 cc. of urine are treated with milk of lime, and the resulting precipitate is removed by filtration, and treated with sulphuric acid and alcohol in a test-glass. Sulphuric acid is added until the solution has an acid reaction. It is then boiled, when the precipitate is decolorised, if bile pigment is present, and the liquid assumes a green tint. With similar treatment urine abounding in indican deposits a bluish-grey precipitate at the outset, but with the subsequent process not a green, but, if any, a yellow or reddish colour develops. Urine which contains haemato porphyrin, on the other hand, is made in this way to exhibit a deep rose colour (see this Chapter : *Hæmatoporphyrinuria*).

4. *Iodine test*: *Kathrein*⁴⁰¹ adds to the urine when freshly passed —otherwise it must be warmed—5–10 drops of a 1 in 10 tincture of iodine. A well-defined green colour marks the presence of bile pigments. *Rosin*⁴⁰² employs the official tincture diluted with 10 per cent. of alcohol. This he pours upon the surface of the urine. A green ring forms in this case.

It may be mentioned here that *Zeehuisen*⁴⁰³ advises that the urine be always diluted to a sp. gr. 1.005 before testing for bile pigments or for albumin or sugar.

*Ehrlich*⁴⁰⁴ has devised the following method for the detection of bilirubin in the urine. The latter is mixed with its own bulk of dilute acetic acid, and, drop by drop, is added a mixture consisting of 1 grm. sulphanilic acid, 15 cc. hydrochloric acid, and 0.1 grm. nitrite of sodium to the litre. A dark discolouration takes place, which, on the further

addition of glacial acetic or other acid, passes to a characteristic violet colour (bilirubin).

C. le Nobel⁴⁰⁵ recommends that the urine be mixed with zinc chloride and a few drops of the tincture of iodine; this produces dichroism. It should be displayed by the urine of jaundice, even if all other reactions fail.⁴⁰⁶

IV. Urobilinuria.—Jaffé⁴⁰⁷ was the first to discover urobilin in the urine. It seldom exists pre-formed⁴⁰⁸ in the recently passed healthy fluid, but the latter holds a chromogen (see *supra*) which yields urobilin on the addition of acid.

According to MacMunn,⁴⁰⁹ the urobilin of febrile urine is different from that of health.

[Febrile or "pathological" urobilin (*MacMunn*), like normal urobilin, is a derivative both of bile and of blood pigment. It is identical with or closely resembles stercobilin. It has a distinct spectrum, and, chemically, is a less highly oxidised product than normal urobilin (see also Chapter I.). The spectrum of pathological urobilin show three bands, one before *D*, one between *D* and *E*, and one at *F*.]⁴¹⁰

This body is present in large amount in the urine of certain morbid states, amongst which may be mentioned fever, and the various affections which are characterised by the disintegration of red blood-corpuscles, as scurvy (*Kretschy*⁴¹¹), and in Addison's disease (*Kummer*⁴¹²).

The increased elimination of urobilin is, however, not constant in Addison's disease.

In two cases investigated by the author urobilin was absent; in another, now under observation, and probably a case of this disease, there is a great increase of urobilin.

[*Mott*⁴¹³ and *W. Hunter*⁴¹⁴ have observed urobilin in the urine in pernicious anaemia, and *Hunter* regards this as a point of much weight in diagnosis. It is also of interest as bearing upon the origin of urobilin. The appearance of urobilin in the urine is associated with an excessive elimination of bile into the intestine (*Hunter*), and with evidence of increased destruction of corpuscles in the portal vein. At the same time, the liver-cells are found to be overloaded with iron. From these facts and others *Mott* concludes that haemoglobin is acted upon by the liver-cells to form urobilin or an allied pigment, which is then excreted by the kidney, while iron accumulates in the liver. *Hunter* assigns reasons for the view that other organs besides the liver effect this decomposition. In one of his cases the substance found in the urine was pathological urobilin.] The above observations, quoted by *Cagney* with reference to the occurrence of large quantities of urobilin in pernicious anaemia, are confirmed by the author's personal experience.

It often happens in cases which would be described as slight jaundice, that a very dark-coloured urine is passed, and this is found on examination to be free from bile pigments, but abounding in urobilin (*Gubler*, *Gerhardt*).⁴¹⁵ This so-called "urobilin jaundice" occurs in connection with hepatic disease, most frequently with hepatic cirrhosis and conges-

tion (*Hayem*).⁴¹⁶ In twelve cases of atrophic and hypertrophic cirrhosis the author has never failed to detect urobilin, and it is, doubtless, valuable evidence of hepatic disease when supported by other symptoms, and in the absence of causes which are known equally to produce urobilinuria. *Rossbach*⁴¹⁷ has observed the latter in a case of multiple neuritis, *Hunter*⁴¹⁸ in pernicious anaemia, *Falcone*⁴¹⁹ in tetanus, and it follows inoculation with Koch's tuberculin. Protracted chloroform narcosis is also recognised as a cause (*Carallero, Kast, and Mester*).⁴²⁰ *Grimm*⁴²¹ found urobilin plentifully in the later stages of digestion.

It is further a fact of great clinical interest that the excretion of considerable quantities of urobilin has been observed to attend on intracranial haemorrhages [*Bergmann, Kunkel*⁴²²], haemorrhagic infarction, retro-uterine haematocele, and extra-uterine pregnancy (*Dick*).⁴²³

These facts are of some weight in diagnosis. In one instance where the clinical symptoms pointed to a serious cerebral disorder, the detection of urobilinuria induced the author to suspect haemorrhagic pachymeningitis, and the autopsy subsequently confirmed the inference.

Apart from its occasional association with liver complaints, the author's experience has been that urobilinuria occurs most commonly in the course of extensive cutaneous haemorrhages due to scurvy, carcinoma, the haemorrhagic diathesis, &c.; extravasation under such conditions was constantly followed by the appearance of urobilin in the urine, and this became more marked as the process fell into abeyance, thus suggesting the inference that the blood-colouring matter discharged into the cuticular tissues was again absorbed and eliminated by the kidneys in the form of urobilin.⁴²⁴ [Cases of this kind which occurred under Dr. Ringer's care are reported by *MacMunn*.⁴²⁵] The individuals in whom this process was going on commonly exhibited a pronounced yellowness of the skin. In such of these cases as afforded an opportunity for post-mortem examination, the bile-duets were invariably found unimpeded, and during life the urine was free from biliary pigments. In the cases of urobilinuria which have come under the author's notice, the skin was very commonly tinged yellow (jaundiced), but this was not without exception, and in such cases he has invariably found bile pigment in the blood (see Chapter I.). From these facts it appears probable that the blood-colouring matter, having been converted into bile pigment, re-enters the circulation as such, and is excreted as urobilin. Since there is no urobilin in the blood, a "urobilin jaundice" cannot be assumed; but in some cases bile pigments derived from the blood are eliminated as urobilin, and in others bilirubin formed in the liver for some reason enters the blood, and is similarly disposed of. [The researches of *Mott* and *Hunter*, noticed above, point to the liver as the place where normal urobilin is sometimes formed.]

Recent investigations by *Leube*⁴²⁶ have made it probable that under some circumstances bilirubin is reduced to urobilin in the kidney.

Urine which contains much urobilin is characterised by its very dark colour; but this does not suffice to distinguish it, inasmuch as other substances, and notably an abundance of the antecedent of indigo, will impart the same quality to the water. Moreover, it will sometimes furnish a beautiful yellow froth like that of jaundice. Such urine has the property of displaying fluorescence in presence of ammonia and zinc chloride. For the detection of urobilin *Gerhardt*⁴²⁷ suggests that a chloroform extract of the urine should be treated with solution of iodine, and caustic potash added, when a beautiful green fluorescence develops.

The test adopted by the author,⁴²⁸ and recommended in previous editions of this work, has been shown by fuller experience not to be serviceable, since the reaction is obtained with hæmatoporphyrin and uroërythrin more readily than with urobilin.

The most accurate qualitative test for urobilin is that employed by *Gerhardt* and *Müller*⁴²⁹ for the estimation of its quantity. A good rough method is that of *Nencki* and *Rotschy*, *Riva* and *Zoja*,⁴³⁰ who extract the urine with amylic alcohol. The test as used by the author is conducted thus:—Fifty cc. of the urine is shaken up with amylic alcohol on a separation-filter, the urine allowed to flow off after some hours, when the tap of the filter is closed and the alcohol pipetted out. The amylic alcohol solution, which is more or less dark in colour, is now treated with concentrated ammoniacal alcoholic solution of zinc chloride. If urobilin be present, the fluid assumes a beautiful fluorescence, and with the spectroscope exhibits the absorption-bands shown in fig. 138.

Riva and *Zoja*⁴³¹ have shown that the red precipitate formed in the amylic alcohol extract by the addition of zinc chloride, contains hæmatoporphyrin (*vide infra*), and this is undoubtedly true; but it is not to be supposed that hæmatoporphyrin and urobilin occur together in *all* urines.

The test with zinc chloride may be applied directly, a few drops of the concentrated ammoniacal alcoholic solution being added to the urine. The resulting precipitate settles, and if the urine be very rich in urobilin the clear fluid fluoresces. Occasionally this and all other tests fail, and resort must then be had to the spectroscope.

Its spectroscopic characters are remarkable. When acid, if rich in urobilin, it displays a distinct absorption-band in the green and blue between Fraunhofer's lines *b* and *F* (fig. 137), and usually extending with diminished intensity beyond *F*. When alkaline, a less well-marked band is seen midway between *b* and *F* (fig. 138). The quantitative estimation of urobilin may be effected by means of *Vierordt's*⁴³² or *Hufner's* spectrophotometer. *Gerhardt* and *Müller's*⁴³³ process is as

follows:—To 100 cc. of urine are added 30 cc. of a baryta mixture containing one part of saturated barium chloride solution and two parts of saturated barium hydrate solution. The precipitate is removed, and one-half—amounting to 65 cc.—of the filtrate is taken, or more if the urine is of low specific gravity. If, on the other hand, it is highly concentrated, it may be diluted first to double its bulk. The fluid is freed from baryta with sulphate of sodium; any urobilin adhering to the solution of barium sulphate is collected by washing the latter with weakly alkaline water, and the filtrate is made slightly acid with sulphuric acid and thoroughly saturated with ammonium sulphate. The ammonium sulphate precipitate is washed on a filter with ammonium sulphate

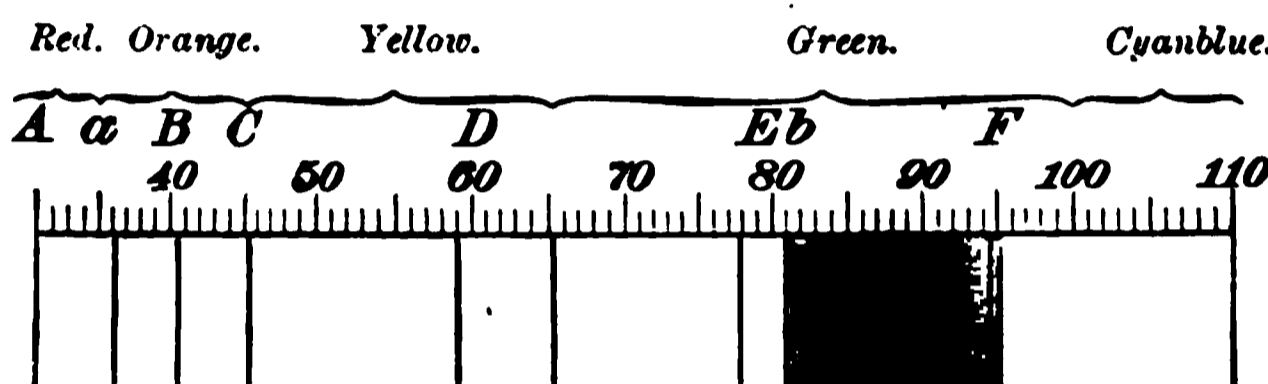


FIG. 137.—Spectrum of Urobilin in Acid Urine.

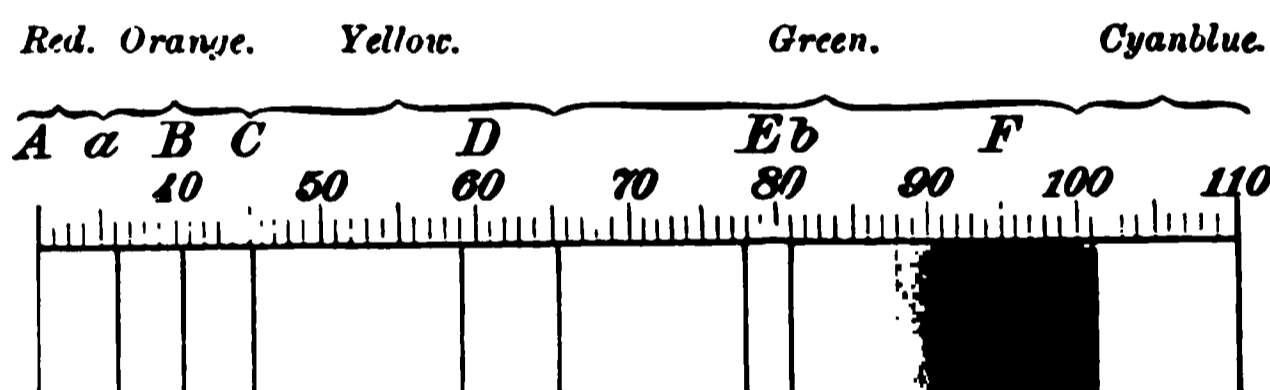


FIG. 138.—Spectrum of Urobilin in Alkaline Urine.

solution (saturated), and together with the filter is placed in a flask and extracted with alcohol acidulated with sulphuric acid (or better, with æther, alcohol, 1 : 2) until the solution ceases to colour. The precipitate need not be entirely dry, but excess of water can be removed by expressing from the filter on absorbent paper. The extracts must now be collected. Urobilin thrown out by saturation with ammonium sulphate, and deposited on the vessel used, is dissolved in a little alcohol, and this is added. The quantity of alkaline solution of urobilin is measured, the percentage of urobilin estimated by the spectrophotometer, and the absolute quantity calculated from this.

For the detection of urobilin *Jaffe's* method⁴³⁴ or *Mehu's*⁴³⁵ (see Chapter VI.), or that of *Riva* and *Zoja*⁴³⁶ just given, may be employed.

V. Hæmatoporphyrinuria.—Hæmatoporphyrin is normally present in the urine (*Garrod*), but only as a trace. Its presence in the urine of diseased states has been noticed [by *MacMunn*, *Le Nobel*, and *E. Salkowski*,⁴³⁷] and cases previously reported by others, and amongst them by *Stokvis* and *Quincke*,⁴³⁸ would appear to be instances of this condition. [(Uro)hæmatoporphyrin, like urobilin, may exist in the urine partly in the form of a chromogen, which becomes a pigment on oxidation (*Halliburton*).⁴³⁹] Viewed by reflected light the urine is opaque and almost black—or in a thin layer, brownish red. The colour is unchanged by boiling. It is, or may be, free from albumin, since hæmatoporphyrin does not contain albumin. When diluted and treated with HCl the characteristic spectrum (*Hoppe-Seyler*⁴⁴⁰) may be seen. This exhibits four absorption-bands, viz., two, narrow and faint, situated, one between *C* and *D*, the second between *D* and *E* but nearer to *E*, and two broad and dark bands, of which one overlaps *D* to the red end of the spectrum, while the other lies between *b* and *F*. These latter may alone be visible, and are alone evidence of hæmatoporphyrin. To detect this substance chemically, about 30 cc. of the urine are taken and treated with alkaline solution of barium chloride; the mixture is filtered, and the precipitate washed first with water and afterwards with absolute alcohol; the precipitate while still wet is rubbed up in a mortar with alcohol and hydrochloric acid, allowed to stand for a while, and then heated in the water-bath. The solution thus obtained, if hæmatoporphyrin be present, should have a reddish colour, and when filtered and examined with the spectroscope, the filtrate should give the two absorption-bands of hæmatoporphyrin (*Salkowski*). This test is neither accurate nor trustworthy. A better is that of *Riva* and *Zoja*.⁴⁴¹ The reddish precipitate obtained by alcoholic ammoniacal zinc chloride in the amylic alcohol extract described above contains hæmatoporphyrin. If this precipitate be suspended in absolute alcohol, the spectrum of hæmatoporphyrin may be obtained from it; or the addition of acids or alkalies will take up the colour from the precipitate. When acid, alcoholic and watery solutions will show violet and exhibit the absorption-bands of hæmatoporphyrin in acid solution (see above). *Garrod's*⁴⁴² test is more accurate and satisfactory than either of these, but complicated, and therefore most suitable when hæmatoporphyrin is very scantily present, and its detection is difficult. *Riva* and *Zoja's* method, on the other hand, will be adopted when the object is to demonstrate an increase in the quantity of this body, a deviation of greater clinical interest. The clinical significance of this state is not yet fully known. [*MacMunn*⁴⁴³ and *Le Nobel*⁴⁴⁴ have described it in Addison's disease, acute rheumatism, pneumonia, measles, pericarditis, typhoid, meningitis, and other diseases.] In some people it follows the use of sulphonal,⁴⁴⁵

and, according to *Herling* and *Schultze*,⁴⁴⁶ is determined also by the administration in excess of trional and tetroonal. It has been observed also in typhoid (*G. Sobernheim*⁴⁴⁷) independently of intestinal haemorrhage (*v. Jaksch*), and it seems likely that a transitory haemato porphyriuria is not uncommon in enteric fever. *Stokvis*⁴⁴⁸ has shown that the condition may result from the absorption of blood from the alimentary canal and its subsequent elimination.

[Haemato porphyrin derived from the blood by the action of sulphuric acid and (uro)haemato porphyrin derived from urine differ in their spectra. *MacMunn* has recently found a pigment, which he regards as intermediate between the two, in three specimens of urine. These urines were of a deep Burgundy-red colour, free from albumin, and on the addition of H_2SO_4 showed the spectrum of acid haemato porphyrin.⁴⁴⁹]

VI. Æther-Sulphuric Acids and their Derivatives (*Indigo-Blue, Indigo-Red, Skatol, Phenol, Parakresol, Pyrocatechin, Hydrochinon*) and the Aromatic Oxy-Acids.

(a.) **Indicanuria.**—Indigo-blue (indigo, indigotin) as such, is rarely present in the urine, usually in decomposed urine, and hardly ever so plentifully as to impart its colour to the fluid. It may always, however, be obtained from urine as a product of the decomposition of salts of indoxyl-sulphuric acid (potassium indoxyl-sulphate).⁴⁵⁰

Indol, a regular product of bacterial putrefaction of albumin (see Chapter VI), is the basis of indican (indoxyl-sulphuric acid).⁴⁵¹ It is oxidised to indoxyl within the system, and by combination with the sulphuric acid present forms indoxyl-sulphuric acid. The decomposition of indoxyl-sulphuric acid yields, besides indigo-blue, other substances of similar character, such as indirubin; but to these no practical interest attaches in the present state of our knowledge.⁴⁵²

With reference to the clinical import of indicanuria, it must be borne in mind that the quantity of indoxyl-sulphuric acid formed varies in health with the food ingested, and it is increased especially by animal diet.

Apart from this, an undue proportion of indican in the urine is a fact of pathological interest, and there are certain diseases in which indoxyl-sulphuric acid is regularly produced in excess.

It was formerly believed that starvation and wasting diseases were attended with the separation of indican (*Senator*⁴⁵³); but more recent observation⁴⁵⁴ has shown that this is dependent on the fact that in such diseases albuminous putrefaction takes place in the alimentary canal, and in consequence there is an increased production of indol, the antecedent of indican. The presence of indican in the urine is very often a sign of intestinal putrefaction, and its quantity in certain cases varies

with the activity of that process.⁴⁵⁵ It may also accompany the decomposition of albumin in other cavities. Thus, in a case of pleurisy with abundant unhealthy exudation, the author has found a profusion of indican in the urine, and when this manifestation arises in the course of peritonitis, it may be taken as an evidence of the character of the disease and of the formation of putrid pus. [Hochsinger⁴⁵⁶ has recently studied this subject in connection with infants. He found that the urine of new-born children was free from indican, and in healthy infants it occurs only in traces. It becomes more abundant in intestinal disorders, and is always most so when these are attended by acute diarrhoea.⁴⁵⁷ Tuberculosis, whether affecting the intestinal tract or not, was always accompanied by profuse indicanuria. Hochsinger attributes the condition to decomposition of milk-albumin in the intestinal tract.] Gehlig⁴⁵⁸ confirms Hochsinger's statements. Singer⁴⁵⁹ detected indican in considerable quantities in the urine of urticaria and other skin affections.

Large doses of thymol were followed by an increased production of indican (*Bohland*). Thymol appears in the urine as thymol sulphuric, thymol glycuronic, thymolhydrochinon sulphuric acids, and as the chromogen of a green pigment (*Blum*). The pigment discovered by *Bohland* is not indican but a derivative of thymol (*Blum*).⁴⁶⁰

In general, therefore, the appearance in the urine of large quantities of indican implies that abundant albuminous putrefaction is progressing actively in some part of the system. Caution must be observed in further narrowing the inference, as to diagnose a gangrenous suppuration, for instance, since in simple constipation a notable indicanuria will very often arise.

*Beckmann's*⁴⁶¹ statement that the intestine is the only source of indican is not to be endorsed. Fœtid pus, though not simple suppuration, and gangrenous processes, lead to its formation on a large scale.

It may be remarked here that the deep-brown colour which usually belongs to urine rich in indican is not due directly to the presence of indoxyl-sulphuric acid, but depends upon the higher oxidation products of indol which accompany it. These bodies bear to indoxyl-sulphuric acid the same relation that the brown, green, or black colouring matters of carbolic urine do to phenol-sulphuric acid.

Detection of Indican.—The methods employed for this purpose proceed upon the principle of splitting up the indoxyl-sulphates of the urine, and obtaining from them a coloured product—indigo-blue.

*Jaffé's Test.*⁴⁶²—A few cc. of the urine are treated with an equal quantity of hydrochloric acid, and, drop by drop, a solution of some hypochlorite is added by means of a glass pipette, and shaken up with the fluid. The chromogen formed by decomposition of indoxyl sulphuric

acid is oxidised into indigo-blue. Care must be taken that hypochlorites are not in excess, since this would alter and bleach the indigo-blue. *Stokvis*⁴⁶³ recommends the admixture of a little chloroform in the process, with the object of dissolving the indigo-blue as it forms. The chloroform solution then takes a blue colour [and the colouring matter is obtained as a deposit after evaporation. Albumin if present should be removed before performing this test, since it forms a blue colour with hydrochloric acid (*Halliburton*).⁴⁶⁴]

*Obermayer*⁴⁶⁵ has suggested a useful modification of *Jaffé's* test. The urine is treated with 1 in 5 solution of sugar of lead, which, however, must not be in great excess, filtered through dry paper, the filtrate mixed with an equal bulk of fuming HCl containing 1 to 2 parts of ferric chloride solution in 500, and then thoroughly shaken for one to two minutes. The indigo-blue formed is then taken up with chloroform.

According to observations made by *Ruzicka*⁴⁶⁵ in the author's clinic, the modification of the *Jaffé* test, proposed by *Asmann*, is unreliable.

*Weber's Test.*⁴⁶⁶—Thirty cc. of urine are mixed with an equal quantity of hydrochloric acid, 1-3 drops of dilute nitric acid added, and the mixture boiled. The fluid assumes a dark colour. If allowed to cool and then shaken up with æther, the presence of indigo-blue is shown by the formation of a blue froth on the surface, while the æther exhibits a rose or violet tint. [*MacMunn* uses chloroform instead of æther, and examines the violet fluid with the spectroscope, which shows an absorption-band before *D* (indigo-blue), and another after *D* (indigo-red). This method is preferable to *Jaffé's* for the detection of small quantities of indigo, which are destroyed by the hypochlorite (*Halliburton*).⁴⁶⁷]

Quantitative Estimation.—The methods of *Jaffé* and *Salkowsky* are the most useful. Their principle is the same as that of the tests for the presence of indican.

*Salkowsky's*⁴⁶⁸ *Colorimetric Process* is perhaps the best. A rough analysis is first effected by determining the quantity of chlorinated lime solution with which indigo forms in greatest abundance. If in this way it is found that the urine contains much indican, 2.5-5 cc. are diluted with water to 10 cc., while, if there be but little indican present, 10 cc. of undiluted urine are taken as the basis of the experiment. In either case an equal quantity of hydrochloric acid is added, and that proportion of chlorinated lime solution which was found in the preliminary reaction to be required. The mixture is then neutralised with caustic soda, and carbonate of soda added to make it alkaline. The indigo-blue which forms is collected in a filter, and then washed with water until it no longer exhibits an alkaline reaction, when it is dried and repeatedly extracted by heating with chloroform until the latter ceases to colour

with it. The chloroform extract is then, by the addition of chloroform, made up to a quantity expressed by a round number of cc., placed in a glass vessel with parallel sides, and the intensity of its colour compared with that of a freshly prepared chloroform solution of indigo-blue of known strength.⁴⁶⁹ To one or other of these, as required, more chloroform is added, until their tint is adjudged equal. From the known constitution of the standard the quantity of indigo-blue derived from the urine (2.5–5 or 10 cc.) taken may be also known, and its percentage may be readily calculated.

From the urine passed in twenty-four hours under ordinary conditions of diet 5–20 mgrms. of indigo-blue can be obtained on an average.

Indigo-Red.—There is now no doubt (*Rosin*⁴⁷⁰) that indigo-red (indirubin) as well as indigo-blue occurs in the urine. It is formed together with indigo-blue when a urine rich in indican is boiled with nitric acid (*O. Rosenbach's* test). For its detection *Rosin* renders the urine alkaline with sodium carbonate, and then extracts the indigo with æther. The inferences which *Rosin* has drawn from the presence of this body in the urine are questioned by other observers, and *Rosenbach's*⁴⁷¹ test cannot be taken as evidence of anything except that the urine contains abundance of the indigo antecedent.

There are certain other aromatic derivatives of the urine which will engage our attention here, both because they are chemically allied to indoxyl-sulphuric acid, and also because pathologically their production is apt to coincide with the manifestation of that substance.

(b.) **Skatoxyl-Sulphuric Acid.**—This body results from the skatol of the faeces (*Brieger*).⁴⁷² It is assumed that skatol, by a process analogous to that undergone by indol, is oxidised to skatoxyl within the body, appearing in the urine as skatoxyl-sulphuric acid. It is probable that the red colour which develops in the urine in the presence of acids is to some extent due to decomposition products of this substance.⁴⁷³

(c.) **Parakresol- and Phenol-Æther-Sulphuric Acid.**—The other members of the aromatic group which occur in human urine in combination with sulphuric acid are phenol (carbolic acid), parakresol, pyrocatechin, and hydrochinon. To the latter we shall have need to refer again. The methods by which these bodies may be detected are highly interesting and clinically instructive.

*Salkowski*⁴⁷⁴ has shown that the urine of patients suffering from ileus and peritonitis, in addition to a large percentage of indican, contains also a considerable proportion of phenol-forming substance, and *Brieger's*⁴⁷⁵ experiments have proved that the elimination of the antecedent of indigo (indoxyl-sulphuric acid) on the one hand, and that of the phenol-producing substances (phenol, parakresol-æther-sulphuric acid) with the aromatic oxy-acids on the other, bear to one another no

constant relation as to activity; and that author found that in diphtheria, scarlatina, and facial erysipelas, phenol was formed in greatly increased quantity, whilst in typhoid, relapsing and intermittent fevers, small-pox, and meningitis, it could be obtained but very sparingly from the urine. These statements are borne out by others.⁴⁷⁶ *G. Hoppe-Seyler*⁴⁷⁷ has reported an increased elimination of æther-sulphuric acid in cholera.

Again, in all cases where albuminous putrefaction is actively progressing in the intestine or other organs, in addition to the salts of indoxyl-sulphuric acid, phenol is increased in the urine; and in general, together with phenol, the other members of the aromatic group become evident in connection with pulmonary gangrene, putrid bronchitis, fœtid pleuritic exudation, and decomposition generally throughout the body.

Detection of Æther-Sulphuric Acids.—For this purpose the urine is first treated with barium chloride in excess to precipitate simple sulphuric acid, and then boiled with hydrochloric acid. If æther-sulphuric acid be present, it is decomposed with the formation of the uncombined acid. This combines with the barium present to form sulphate of barium, and a white precipitate is deposited.

The quantitative estimation of the phenols (phenol, parakresol) is conducted in the manner described below, and their presence may be determined by the tests given in Chapters V. and VI. It must be mentioned, however, that the investigations of *Rumpf*⁴⁷⁸ have shown that an accurate quantitative analysis cannot be made by any of the methods hitherto in use. On the other hand, *Kossler* and *Penny's*⁴⁷⁹ process gives good results, as is shown by *Strasser's*⁴⁸⁰ observations in the author's clinic. For the comparative investigation of these substances the original work of *Brieger*⁴⁸¹ may be consulted.

Quantitative Estimation of Æther-Sulphuric Acids.—The percentage of this body in the urine may be best determined by *Baumann's*⁴⁸² method as modified by *Salkowski*.⁴⁸³

To 200 cc. of urine is added a like quantity of alkaline barium chloride solution (two parts saturated solution of baryta, and one part of solution of chloride of barium saturated in the cold). The mixture is allowed to stand for some minutes, and then passed through a thick filter which has been carefully dried. Of the filtrate, *which must be perfectly clear*, 100 cc. are taken and rendered strongly acid with 10 cc. of hydrochloric acid (sp. gr. 1.12), then boiled and heated on the water-bath until all the precipitate which forms has settled. The beaker may be heated on an iron plate coated with asbestos, and allowed to remain till its contents are cold. The precipitate is next collected and placed on a filter of Swedish paper which has previously been washed with dilute hydrochloric acid, and care must be taken that the filter is not allowed to empty itself entirely during the process. With the aid of a

glass rod protected with a ring of india-rubber, and rinsing with boiling water, the entire precipitate is placed on the filter, and is there washed with boiling water until the filtrate which passes through fails to give a precipitate with dilute sulphuric acid, thus showing the absence of free chloride of barium. Should it happen that the fluid passing through the filter is turbid, this may be due to the presence of soluble substances, as phenols produced by the decomposition of the compound acids. To ascertain their character in this case, the cloudy filtrate is placed in a beaker on the water-bath heated to boiling-point, when, if the turbidity be due to phenols, these pass off in vapour and leave the fluid clear. Where, on the other hand, the appearance is caused by the barium precipitate having passed through the filter, the turbidity will not be removed in this way, and the experiment is spoiled. The precipitate is next washed with boiling alcohol, and finally with æther, and together with the filter-paper placed upon a platinum crucible of known weight and heated for a long time. After this the platinum crucible is raised to a white heat, allowed to cool, and then weighed again. The calculation is made as follows:—233 parts by weight of sulphate of barium correspond to 98 parts by weight of sulphuric acid (H_2SO_4), and consequently the quantity of sulphuric acid in 100 cc. of the urine may be computed by the formula—

$$x = \frac{98}{233} \times M = 0.4206 \times M$$

Where

x = the quantity of sulphuric acid required.
 M = the quantity of barium sulphate found.

If the object be to determine the total quantity of the sulphuric acids (simple and compound), so as to find out their respective proportions in the urine, the latter is filtered. Another 100 cc. are taken and treated with 10 cc. of hydrochloric acid (sp. gr. 1.12), then boiled for fifteen minutes, and chloride of barium solution added in excess. The remainder of the process is that described above. The difference between the total quantity and that of æther-sulphuric acids obtained expresses the proportion of simple sulphuric acid in the urine.

For the methods of estimating sulphur when present in the urine in other forms see pp. 377-403.

Quantitative Estimation of Phenols.—The phenols (phenol and para-kresol) which have passed over in the distillation of a known quantity of urine previously acidulated are estimated in the form of tribromo-phenol by Landolt's method, with the precautions suggested by Baumann and Brieger.⁴⁸⁴

One-fourth of the urine passed in twenty-four hours is mixed with

one-fifth its bulk of hydrochloric acid and distilled. Distillation is continued until the distillate ceases to colour with bromine water, after which it is filtered. All the fluid which has passed through—including that tested during the process—is now treated with bromine water until a permanent yellow colour is attained. The precipitate is allowed to settle for two or three days, when it is separated on a filter which has been weighed and carefully dried over sulphuric acid; it is then washed with bromine water and dried over sulphuric acid in the dark until it has acquired an approximately constant density. It is then weighed on the filter, and the difference between the result and the weight of the filter recorded expresses that of the tribromo-phenol formed. From this the quantity of carbol in the urine may be estimated thus:—331 parts by weight of tribromo-phenol correspond to 94 parts by weight of carbol, and the following formula results:—

$$x = \frac{94}{331} \times M = 0.2839 \times M$$

Where

x = the quantity of carbol required.

M = the quantity of tribromo-phenol found.

This method may also be employed for the analysis of vomited matters in cases of carbolic acid poisoning (compare Chapter IV.).

Another method.—The following admirable and very simple method is based on the analytical process of *Koppeschaar, Messinger, and Vortmann*:⁴⁸⁵—500 cc. of urine are rendered feebly alkaline, and evaporated down to 100 cc., the fluid received in a distillation flask, 25 cc. of concentrated sulphuric acid added, the fluid distilled, water added after distillation, and the process repeated several times. The distillates first obtained are collected; the later ones are better treated separately. To each is added calcium carbonate to neutralise the fluid, when it is again distilled. The distillate, or an aliquot part of it, is placed in a flask with a ground-glass stopper, treated with deci-normal solution of soda till it is strongly alkaline, and then the flask securely stoppered and immersed in boiling water for a long time. To the fluid, while still hot, is added 15–25 cc. of deci-normal iodine solution, and as much more as corresponds to the quantity of deci-normal soda solution previously added; the flask is again stoppered and well shaken. The fluid should have a brown tint. After cooling it is made acid, and the resulting free iodine titrated with deci-normal thio-sulphate solution. The process is repeated with each of the original distillates. Each cc. of the iodine solution used represents 1.567 mgm. of phenol or 1.8018 mgm. of para-kresol, and the total quantity of the iodine solution used multiplied by the figure 1.567 or 1.8018 gives the amount of phenol or of kresol

respectively in the urine expressed in mgrms. It is better to estimate the phenols as kresols.

The amount of phenols computed as kresols eliminated in the urine daily on a mixed diet is 0.081–0.122 grm., or, according to *Strasser*,⁴⁸⁶ 0.06–0.08 grm. The latter obtained as the maximum quantity in a case of gangrene of the foot 0.40 grm. estimated as kresol.

(d.) **Pyrocatechin.**—This body, like the others, occurs in the urine only in combination with sulphuric acid ; and it is, if not an invariable, at all events a very frequent constituent of that fluid.⁴⁸⁷ The urine containing it is characterised by the fact that it is colourless when passed, but becomes dark after exposure to the air ; and the change is expedited by the addition of caustic potash. When boiled with hydrochloric acid, it becomes a powerful reducing agent. An ammoniacal solution of silver exposed to it will quickly deposit metallic silver in the cold.

These properties, however, are not sufficient to determine the presence of pyrocatechin. To do this it must be isolated, and the following method is the best for the purpose (*Baumann*).⁴⁸⁸ The urine is made strongly acid with hydrochloric acid, heated on the water-bath, and, when cool, extracted with æther. The æthereal extract is shaken up with soda solution until it ceases to turn yellow. When the æther has been driven off, the fluid remaining is extracted with a small quantity of saturated solution of sodium sulphate, water added, and the solution distilled until phenol ceases to pass over (see p. 348). The residue is extracted with æther, and the latter evaporated away. The remaining fluid is of a syrupy consistence and crystalline from the presence of pyrocatechin ; to this, diluted with water, is added a little sugar of lead, care being taken that the salt is not in excess. The sugar of lead precipitate, which contains all the pyrocatechin, is treated with sulphuric acid and extracted with æther. Pyrocatechin passes over in the æthereal extract, and when the æther is distilled off, it remains as a more or less pure crystalline substance. If the character of the latter is not sufficiently apparent, it may be recrystallised from benzol in the form of prisms belonging to the tetragonal system. If some of these be dissolved in water in a watch-glass and a few drops of a very dilute solution of perchloride of iron be added, an emerald-green colour develops, and this changes to violet on the addition of a little ammonia.⁴⁸⁹

(e.) **Hydrochinon.**—It has been ascertained by *Baumann* and *Preusse*⁴⁹⁰ that hydrochinon appears in the urine after carbolic acid poisoning, and these authors believe that to its presence the dark colour of that fluid after the exhibition of carbolic acid is due. It is always in the form of æther-sulphuric acid in the urine, and the process for its detection is the same as that for pyrocatechin.⁴⁹¹ The filtrate, after the addition of sugar of lead, contains hydrochinon (*r. supra*). Sulphuric

acid is added, and then carbonate of baryta; the fluid is heated, filtered, and the filtrate extracted with æther. When the latter is driven off, crystals of hydrochinon remain.

The crystals of hydrochinon belong to the rhombic system, and they crystallise readily from their solution in toluol.

According to *Baumann* and *Preusse*,⁴⁹² when rapidly heated in an open test-tube, hydrochinon forms violet fumes, which condense as an indigo-blue sublimate, and in the application of this property we possess a very sufficient test for its presence. When boiled with ferric chloride it emits the odour of chinon.

(f.) **The Aromatic Oxy-Acids.**—The aromatic oxy-acids which have been proved to exist in the urine are paroxyphenyl-acetic acid, paroxypropionic (hydroparacumaric) acid,⁴⁹³ paroxyphenyl glycolic acid,⁴⁹⁴ and oxyamygdalic acid,⁴⁹⁵ to which must be added uroleucic (trioxyphe nyl-propionic) acid (*Kirk*, *Wolkow*, and *Baumann*)⁴⁹⁶ and homogentisic (dioxyphenyl acetic) acid. In view of the significance which the researches of *Wolkow* and *Baumann* have conferred upon these latter substances, there will be need to refer to them separately under the head of *Alkaptonuria*.

Detection of the Aromatic Oxy-Acids.—Twenty cc. of urine are treated with hydrochloric acid, and heated for some time on the water-bath to expel the volatile phenols. The fluid is then allowed to cool, and repeatedly extracted with æther. The æthereal extract is shaken up with a weak solution of carbonate of soda. The oxy-acids are taken up by the latter, whilst the phenols present are retained by the æther. The alkaline solution is now acidulated with sulphuric acid and extracted with æther. The latter is allowed to evaporate, and the residue dissolved in water is submitted to Millon's test. A red coloration shows the presence of aromatic oxy acids.

In a similar manner an approximate quantitative estimation of these bodies may be effected.⁴⁹⁷

VII. Alkaptonuria.—Although the substances with which we have to deal under this heading belong properly to the aromatic series, and are oxy-acids, the special clinical importance which attaches to them makes it convenient to adopt *Baumann's* classification, and treat them apart. By the term alkaptonuria is meant the condition in which the uroleucic acid of *Kirk* and the homogentisic acid of *Wolkow* and *Baumann* occur in the urine. There is no doubt that the urine in certain cases (see above) contains these acids in addition to pyrocatechin. *Berdeker*⁴⁹⁸ gave to a substance resembling these, which he discovered in the urine, the name of alkapton. A similar substance was discovered by *Ebstein* and *J. Müller*⁴⁹⁹ very abundantly present in a child's urine, and *Fasching* and *Fleischer*⁵⁰⁰ have obtained evidence of such in cases

of phthisis. Interesting particulars have been more recently published by *Garnier* and *Voirin*,⁴⁹⁹ *Geyger*,⁴⁹⁹ and *Embden*.⁵⁰⁰ The urine has characters very similar to those conferred upon it by the presence of pyrocatechin (see p. 350). *Baumann* and *Wolkow* believe that the production of the acids in question is due to an anomalous metabolic process of life-long continuance, by which the tyrosin (paroxyphenyl-amidopropionic acid) of the system is perverted, probably by the action of some definite micro-organism. For the method of recognising homogentisic acid the reader is referred to the very interesting treatise of *Baumann* and *Wolkow*.⁵⁰¹ The condition of the urine known as alkapttonuria may be said to exist when the urine answers the description given at p. 350, thus showing that it contains a large amount of pyrocatechin.

VIII. Inosituria.—Inosite occurs in small quantity in the urine in cases of diabetes insipidus and in albuminuria. For its detection it must be separated from the urine. To this end *Cooper-Lane's*⁵⁰² method is the best. According to *Magneune*,⁵⁰³ inosite is obtained as hexahydrobenzol.

IX. Melanuria.—The urine of persons suffering from pigmented tumours sometimes contains a substance to which the name of melanin has been given, but of which the chemical constitution is not yet sufficiently established. It is sometimes held in solution, and more rarely suspended in the form of small granules. It very seldom happens that the fluid has a dark colour when passed, but it generally blackens intensely when submitted to oxidising agents (sulphuric and hydrochloric acids and ferric chloride). This fact would point to the conclusion that the body in question is not melanin, but a chromogen—melanogen—analogous to that which precedes the formation of urobilin. Such urine turns dark on exposure to air. The pigment can be partially separated from the urine by means of acetate of lead or perchloride of iron. It is insoluble in cold alcohol, æther, and acetic and dilute mineral acids. It is soluble in boiling concentrated mineral acids, in boiling lactic and acetic acids, in concentrated solutions of caustic potash and soda, and in ammonia. It contains iron, sulphur, and nitrogen. The most sensitive test for the presence of melanin is the addition of bromine water (*Zeller*),⁵⁰⁴ which causes the urine to deposit a yellow precipitate, which gradually blackens.

More recent experience has shown that a fairly concentrated solution of perchloride of iron serves well to detect its presence (*v. Jaksch, Pollak*).⁵⁰⁵ A few drops of this reagent will cause the fluid to turn grey; and if more be added, a precipitate of phosphates falls, carrying the colouring matter with it, and again dissolves with an excess of the solution.

Sodium nitro-prusside with caustic potash and acetic acid gives a deep-blue colour (*Thormählen*),⁵⁰⁶ which depends probably on the formation of soluble and insoluble Berlin-blues (*v. Jaksch*).

In connection with this subject the reader may refer to the opinion of *Kruckenbergs*⁵⁰⁷ and *Salkowski*⁵⁰⁸ (see under *Kreatinin*), that in *Weyl's* test for *kreatinin*, boiling with acetic acid similarly produces Berlin-blue.

It would appear that in these cases the blood also contains abundance of pigment (see Chapter I.). In a case of melano-sarcoma of the liver verified by autopsy, this was the condition of the blood, and the urine disclosed the characters given above (*v. Jaksch*).⁵⁰⁹

The latter, however, cannot always be obtained by the action of the nitro-prusside salt on melanin isolated from the urine, and the reaction must *not* be regarded as a test for melanuria, or only when other tests (and especially that with perchloride of iron) have shown the presence of melanin or melanogen. Moreover, the Berlin-blue reaction can be obtained in urine which is *free from melanin*. Thus, in the case of children suffering from prolonged constipation, it has been had at the same time that the fluid was rich in acetone or diacetic acid and indoxyl-sulphuric acid (*v. Jaksch*), and in a case of diabetes by *Dreschfeld*,⁵¹⁰ when its nature and the presence of the substances just named were probably established. It would appear, therefore, that in these conditions also a body is present which gives Berlin-blue with nitro-prusside compounds. Possibly this is indol. Investigations with that body derived from a preparation of picrate of indol gave the same result (*v. Jaksch*). The practical significance of this condition is greatly limited by the fact that the urine may contain a large quantity of melanin in wasting diseases, whilst that derived from individuals suffering from melanotic cancer or sarcoma may be entirely free from it. *Senator*⁵¹¹ has recently confirmed this view by a series of clinical observations. Nevertheless, as an adjunct in diagnosis, the tests given are of undoubted utility.⁵¹²

X. Acetonuria.—Normal urine contains traces of acetone (*physiological acetonuria*, *v. Jaksch, de Boeck, A. Sloesse*),⁵¹³ but this body occurs in excessive proportion under certain morbid conditions (*pathological acetonuria*). In association with diseases we may distinguish (1) febrile acetonuria; (2) diabetic acetonuria; (3) acetonuria accompanying certain forms of cancer independently of inanition; (4) acetonuria of starvation; (5) the production of acetone in psychoses; (6) acetonuria as an expression of auto-intoxication; (7) acetonuria in derangements of digestion;⁵¹⁴ (8) acetonuria in chloroform narcosis (*Jufé, Becker, Greven*).⁵¹⁵ In this connection the presence of acetone is referred by *Becker, Abram* to albuminous decomposition. The occurrence of acetonuria in association with psychoses has lately been illustrated by *Wagner*⁵¹⁶ with a great profusion of clinical facts. The commonest of

these forms is febrile acetonuria. It belongs to children as well as to adults (*Baginsky*).⁵¹⁷ It does not belong especially to any particular fever. In connection with diabetes the appearance of acetone in the urine indicates an advanced stage of the disease, but does not otherwise affect the prognosis. Of greater consequence⁵¹⁸ are those cases in which much acetone is found in connection with grave symptoms of cerebral irritation, less often of depression. Acetonuria existing alone (auto-intoxication with acetone) tends to a favourable termination (*c. Jaksch*).⁵¹⁹ Finally, it should be noticed that recent researches have shown that an abundance of nitrogenous food tends to the production of acetonuria. The principal source of acetone was shown by the author some years since to be decomposition of proteids both of the body and taken as food. This view has lately had support from *Rosenfeld*.⁵²⁰ It doubtless originates in other ways also, but the inferences recently drawn from his observations by *Hirschfeld*⁵²¹ cannot be readily accepted.

Detection of Acetone.—A rough test for acetone is that of *Legal*. A quantity of the urine (several cc.) is treated with a few drops of a freshly made and somewhat concentrated solution of sodium nitro-prusside, and with a moderately strong solution of caustic soda or potash. The fluid develops a red colour, which rapidly disappears, and if acetone be present, gives place to purple or violet-red on the addition of a little acetic acid. In the absence of acetone the purple-red tint does not form on the addition of acetic acid.

For purposes of greater accuracy it is necessary to distil the urine, and to apply to the distillate the tests presently to be described. To do this, one-half to one litre of the urine may be taken, and a little phosphoric acid may be placed with it in the retort to prevent the evolution of gases.

To avoid the use of acid and still guard against evaporation, the author almost invariably uses the steam-jet. Steam is generated in a tin kettle furnished with a water-gauge and safety-valve, and led into a flask containing the urine. The flask is connected with the kettle and with a distillation apparatus by air-tight tubing. By proceeding thus, errors, as from the formation of aldehyde (*Sulkowski*)⁵²² (see p. 165), are prevented.

Of the distillate 10-30 cc. may be taken and tested with—

(1.) *Lieben's Test*.—To several cc. a few drops of iodo-potassic iodide solution and caustic potash are added. If more than a trace of acetone be present, an abundant precipitate of iodoform crystals is deposited. This test is very reliable, and will serve even for the detection of traces of acetone.

(2.) *Reynolds' Test*.—This depends on the property which acetone possesses of promoting the solution of recently formed mercuric oxide. It is conducted as follows:—The yellow precipitate (mercuric oxide)

obtained by the reaction of mercuric chloride with an alcoholic solution of caustic potash is added to the distillate from the urine, which is then filtered, and to the clear filtrate sulphide of ammonium is cautiously added. If acetone be present, some of the mercuric oxide will have dissolved, and a black ring (sulphide of mercury) forms at the plane of contact with the ammonium sulphide.

Legal's Test, already described, may be applied also to the urinary distillate, but it is less to be relied upon than the others, since parakresol, which also passes over in distillation, exhibits a similar reaction.⁵²³

Quantitative Estimation of Acetone.—This may be effected by *v. Jaksch*'s method as modified by *Nencki*.⁵²⁴ Very good results are obtained by the process first devised for scientific purposes by *Messinger*,⁵²⁵ applied to the investigation of urine by *Huppert*,⁵²⁶ and rendered available for clinical use by *v. Engel* and *Deroto*.⁵²⁷ It is conducted thus:—The urine is first examined by *Legal's* test, and, according to the result, 20–50, or at most 100 cc., are placed in a flask and made up to 100 cc. with distilled water and 2 cc. of a 50 per cent. acetic acid solution. This flask is connected by a long glass tube with the cooler, in front of which is a distillation flask, and in front of that a bullet apparatus filled with urine. Distillation is carried on until $\frac{9}{10}$ ths of the original volume of the fluid have passed over. A portion of the residue is submitted to *Lieben's* test, and if this shows the presence of acetone, the result must be rejected and the process commenced over again after the addition of more distilled water. To the distillate 1 cc. of dilute (1 in 8) sulphuric acid is added and the mixture distilled. The second distillate is poured into a flask of 1 litre capacity fitted with a polished glass stopper, and also, for distillation purposes, with a doubly perforated cork, and having a bullet apparatus full of water in front of it. When distillation is completed, the flask is closed with its glass stopper and the fluid carefully titrated according to *Huppert's* directions with $\frac{1}{16}$ th normal iodide solution and $\frac{1}{15}$ th normal hyposulphite solution. These solutions being used, 1 cc. of the iodo-solution corresponds to 0.967 mgm. of acetone. The researches which *v. Engel* has pursued by this method have greatly extended our previous knowledge concerning the secretion of acetone, and they have also established the facts on a secure basis. *A. Jolles*⁵²⁸ has devised a quantitative estimation method, based on the acetone-phenylhydrazin test. *Parlato*⁵²⁹ employs the vaporimeter for the purpose. *Sapino*⁵³⁰ advises that the iodoform in the urinary distillate be dissolved in aether, converted into sodium iodide, and the latter titrated with silver nitrate. This method should give trustworthy results.

XI. Diaceturia.—By the term diaceturia is meant the condition in which diacetic acid appears in the urine. It is always pathological,⁵³¹ and occurs in diabetes (*Gerhardt*) and fevers (*v. Jaksch*, *Deichmuller*,

Seifert), and also idiopathically as a form of auto-intoxication. It is most common in children as a concomitant of fever,⁵³² and is then generally devoid of serious significance, but in adults it is a symptom of grave import. In febrile and diabetic states the development of diaceturia commonly forebodes the advent of coma.

Urine holding diacetic acid is always rich in acetone, and in presence of perchloride of iron develops a Bordeaux-red. This property, however, does not serve to distinguish the presence of the former body, since it belongs equally to a number of substances which are apt to exist in the urine.⁵³³ For its detection the following process may be adopted.

To the urine a fairly concentrated solution of perchloride of iron is cautiously added, and if a phosphatic precipitate forms, this is removed by filtration and more of the perchloride of iron solution supplied. If the Bordeaux-red colour appears, one portion of the urine is boiled, whilst another is treated with sulphuric acid and extracted with æther. If now the urine which has been boiled shows little or no change, whilst the perchloride of iron reaction in the æthereal extract is no longer evident after 24–48 hours; and if at the same time (on testing the urine directly and its distillate) it is found to be rich in acetone, the condition may be inferred to be that of diaceturia.

*K. H. Mörner*⁵³⁴ adds to the urine to be examined sodium iodide and ferric chloride in excess, and boils. If diacetic acid be present, irritant fumes (iodo-acetone) are generated. The author has found that urine holding acetone, though free from aceto-acetic acid, behaves similarly. This test is therefore worthless.

XII. Lipaciduria.—By this term is meant the condition in which volatile fatty acids are found in the urine (*v. Jaksch*,⁵³⁵ *v. Rokitansky*⁵³⁶). These bodies occur there in traces normally, especially formic, acetic, and butyric acids; and they may be derived from healthy urine in considerable quantity by the use of oxidising agents.⁵³⁷ They are also a product of alkaline fermentation.⁵³⁸

As a manifestation of disease, on the other hand, they are often present in quantity in the simple urine. Thus, in the urine of fevers, of hepatic diseases affecting the proper structure of the liver, and in diabetes, formic, acetic, butyric, and recently also propionic acid have been detected.

There is no special diagnostic significance attaching to this condition; in general, it is determined by the same causes which produce febrile acetonuria.

For the detection of fatty acids the urine is distilled with phosphoric acid, and the distillate carefully neutralised with carbonate of soda, evaporated to dryness on the water-bath, the residue extracted with boiling alcohol, filtered, again evaporated, dissolved in water, and the

solution submitted to the tests mentioned at p. 241. The principal reactions are shortly recapitulated here.

1. A little of the urine is treated with sulphuric acid and alcohol. An odour of acetic æther indicates the presence of acetic acid.
2. To another portion perchloride of iron is added. The specimen assumes a red tint, which disappears on boiling, and a rusty precipitate remains.
3. The addition of nitrate of silver causes a white precipitate, which rapidly blackens if formic acid be present.

With reference to the appearance in the urine of other organic acids, see pp. 346, 351.

XIII. Lipuria.—Small quantities of fat are often seen in the urine of chronic nephritis with a very fatty state of the kidney (see pp. 269, 287), in phosphorus-poisoning,⁵³⁹ and diabetes mellitus. Fat in large proportion was found by *Ebstein*⁵⁴⁰ in a remarkable case of pyonephrosis.

Fat is also a common manifestation in chyluria, and it is a physiological constituent of the urine of pregnant women.

*Schlossmann*⁵⁴¹ observed fat in the urine of children who had been taking castor or olive oil.

Its presence is sufficiently apparent. The urine containing it is usually very turbid, and clears when shaken up with æther. The fatty particles may be separated by means of Stenbeck's sedimentator (see p. 256). It is apt also to hold globules of fat, which are easily recognisable by their powerful refracting properties; and it is not uncommon for this substance to occur in the form of needles, as it does in the faeces (p. 233), especially in connection with chronic nephritis and septicæmia.⁵⁴²

XIV. Chyluria.—By this term is meant the simultaneous appearance at intervals of fat and albumin in the urine, apart from the manifestation of other morbid constituents, such as casts, renal epithelium, &c. The sediment, however, usually contains red and white blood-corpuscles in small numbers.

The urine under these circumstances tends to form coagula of fibrin on standing, and occasionally it gelatinises throughout. Hitherto chyluria has been met with almost exclusively in the tropics, and in persons who have lived there for a long time, and it has been shown by *Wucherer* and *Lewis* (see Chapter I.) to depend upon the invasion of the urinary tract by *Filaria sanguinis hominis*. The embryo of this parasite is generally to be found in the urine; and the chemical investigations of *Grim*⁵⁴³ make it appear that in the majority of cases the abnormal condition of the urine is due to unnatural communications between the lymphatics and urinary passages affected by filaria. The subject, however, needs further elucidation, inasmuch as chyluria is occasionally observed in persons who have never lived in the tropics.⁵⁴⁴

*Langgaard*⁵⁴⁵ has detected large quantities of cholesterol in the urine in a case of chyluria.

XV. Oxaluria.—It has been stated already that oxalic acid occurs in healthy urine; but it is subject to very great increase in certain morbid states, and the condition is then called oxaluria.

Oxalates may remain in solution in the urine, and it is important to be able to determine absolutely the quantity of oxalic acid present as such. This can be done by a modification of *Neubauer's* method (*Furbringer* and *Czapek*).⁵⁴⁵

*Quantitative Estimation of Oxalic Acid, Neubauer's Method.*⁵⁴⁷—The urine passed within twenty-four hours is accurately measured, and treated first with calcium chloride and ammonia, then with acetic acid until it has a slightly acid reaction, and afterwards a little alcoholic solution of thymol is added to restrain the development of micro-organisms. The mixture is allowed to stand for some time, when the white precipitate which forms is separated on a filter, and (together with the latter) is placed in hydrochloric acid, gently heated, the fluid filtered off, and the filter washed with water until it has no longer an acid reaction. The collected filtrate is evaporated to a small bulk in a capsule on the water-bath, then placed in a strong glass cylinder, and the capsule in which it was evaporated is washed with dilute hydrochloric acid and water, the washings being added to the fluid in the cylinder. Ammonia solution is then poured upon the surface of the latter, and the whole is tinted with a few drops of tincture of litmus. The mixture is allowed to stand for a considerable time. The precipitate which has formed is obtained on a so-called ash-free filter, the ash constituent of which has previously been accurately ascertained, and the oxalate (of lime) which adheres to the walls of the cylinder is removed on a glass rod guarded with an india-rubber ring, and added to the precipitate on the filter. The latter is next freed from chlorine by washing with water, and rinsed with acetic acid. The filter is then dried, and ignited on a platinum crucible, which is heated to a constant weight in the blow-pipe flame. By this means oxalate of lime is changed into lime. Now as 56 parts of lime correspond to 90 parts of oxalic acid, the quantity of the former obtained when multiplied by 1.6071 shows the quantity of oxalic acid in the urine taken.⁵⁴⁶

In healthy urine the amount of oxalic acid passed in twenty-four hours is 0.02 gm. (*Furbringer*).

An excess of oxalic acid is occasionally found in diabetes, and especially when the proportion of sugar diminishes (vicarious oxaluria).⁵⁴⁸

Oxaluria is also known as an affection *sui generis* (oxalic acid diathesis, idiopathic oxaluria), (*Cantani*).⁵⁵⁰

It must be admitted that our knowledge of this condition as a clinical symptom is very defective, but the author's experience induces him to adopt the conclusion of *J. Beybie*⁵⁵¹ and *Cantani*, that there are

certain complaints, characterised by pains in the back and loins and attended with rapid emaciation, in which the only objective symptom besides is an excessive elimination of oxalic acid with the urine. *Neiderl*⁵⁵² observed in a complaint with nervous symptoms more than 0.5 grm. of oxalic acid per litre of urine; and *Kisch*⁵⁵³ found in nine cases of extreme lipomatosis only one in which there was an increase of 0.040 grm. in the litre. *Abeles*⁵⁵⁴ has shown that the quantity of oxalic acid eliminated in the urine is not increased by the administration of oxalates with the food.

XVI. Cystinuria.—This is a condition of rare occurrence, and clinically of little importance, since it is only accidentally by the formation of calculi that it gives rise to trouble. It is usually chronic in its course. It should be mentioned that *Ebstein*⁵⁵⁵ has found cystin concurrently with albumin in the urine of acute articular rheumatism (comp. p. 283).

The researches of *Stathagen*, *Brieger*, *v. Udransky*, and *Baumann*⁵⁵⁶ have shown that such urines also contain diamines, and in particular putrescin, cadaverin, and a diamine which is isomeric with the latter. These bodies occur at the same time in the faeces of such patients (see Chapter VI.), while both urine and faeces of healthy persons are free from them. It is possible that they originate in a special form of intestinal infection, are absorbed from the alimentary canal, and eliminated together with cystin in the urine.

XVII. The Uric Acid Diathesis.—Although the deposition of a very abundant sediment of urates in the urine does not warrant the inference that uric acid is excreted in excess, there is no doubt that there are certain processes in the system, the chief evidence of which is such an increase in the elimination of uric acid, and it is important to possess the means of estimating this condition.

Many methods have been devised for this purpose, as those of *Fokker* and *Salkowski*, and in recent years by *Haycraft*,⁵⁵⁷ *Czapek*, and *W. Camerer*.⁵⁵⁸ That of *Fokker*,⁵⁵⁹ as modified by *Salkowski*,⁵⁶⁰ depends upon the comparative insolubility of urate of ammonia. Those of *E. Salkowski*⁵⁶¹ and of *E. Ludwig*⁵⁶² are based on the estimation of the almost insoluble double silver salt of uric acid. *Hopkins'* method more recently introduced excels the others in accuracy and rapidity.

Ludwig's process has the advantage that it can be carried out completely in twelve to fourteen hours, and it is further serviceable as a qualitative test for uric acid in the other secretions, and in the blood as well as in the urine.

For its application the following solutions are needed:—

i. *An Ammoniacal Silver Solution.*—This is prepared by dissolving 26 grms. nitrate of silver in distilled water, and adding ammonia until

the brown precipitate thrown down at first is again dissolved. The fluid is made up to a litre, placed in a well-stoppered flask, and protected from the light.

ii. *A Magnesia Mixture.*—A hundred grms. of crystallised magnesium chloride are dissolved in water, and a large excess of ammonia added, and then ammonium chloride until the precipitate (magnesium hydrate) is entirely dissolved. The fluid so derived should be tolerably clear. It is made up to a litre, and placed in a stoppered bottle till required.



FIG. 139. Ludwig's Filter (actual size).

iii. *Solution of Sodium or Potassium Sulphide.*—Fifteen grms. of caustic potash or 10 grms. of caustic soda are dissolved in a litre of water, and one half of the fluid is saturated with sulphuretted hydrogen, after which the other half is added to it. The potash or soda used must be entirely free from nitrates and nitrites, and to this end it is well to use caustic soda prepared from metallic sodium.

A hundred or 200 cc. of urine are measured off in a dry glass cylinder and carefully poured into a beaker of 200–300 cc. capacity. Ten or 20 cc. (according as 100 or 200 cc. of urine has been taken) each of solutions i. and ii. are mixed together in a measure-glass, and ammonia is slowly added until the precipitate is dissolved. The clear fluid is then poured from the cylinder in which the urine was measured, and added to the latter in the beaker-glass, and the mixture stirred for some time. The precipitate which forms is allowed to stand for half-an-hour or an hour, after which it is placed with the fluid on a filter, and two or three times the quantity of water, to which a little ammonia has been added, is supplied. For this purpose Ludwig employs an aspirator, but it is not necessary, since filtration proceeds rapidly enough without it.

The precipitate and the filter together are placed in the beaker, and 10 or 20 cc. (according to the quantity of urine taken) of solution iii., diluted with an equal quantity of water, is heated to boiling in a flask, added to the precipitate in the beaker, and the mixture frequently stirred, 40 cc. of boiling water supplied, and the mixture heated over a flame until it begins to boil. It is repeatedly stirred while allowed to cool, and passed through a filter, which is afterwards washed two or three times with boiling water and collected in a large

capsule. The filtrate is treated with hydrochloric acid until it has a feebly acid reaction, and is then concentrated on the water-bath to a volume of 10-15 cc. Uric acid begins to separate at this point in crystals, which are often of a beautiful white colour.

The best plan is to continue evaporating, without regard to the quantity of fluid remaining, until the point is reached when uric acid begins to separate from the hot solution.

The fluid is now allowed to cool for an hour, when the separation of uric acid will be completed. The precipitate is brought upon a Ludwig's filter arranged with glass-wool. This instrument consists of a glass tube about 14 cm. long and 2 cm. in diameter in its upper point, growing rapidly narrower below, and constricted to little more than capillary calibre at a point 4 cm. from its lower end (fig. 139). The lower end is cut off obliquely. The tube is packed from the point of constriction upwards as far as the beginning of the broad part with glass-wool, which is best introduced by means of a slender glass rod, and in such a manner that the obstruction is densest below, and less compact as it proceeds upwards.

To facilitate this, the glass-wool with which the funnel is to be blocked may be previously moistened with a little æther.

The author prefers asbestos to glass-wool. It answers the purpose well, and does not irritate the skin.

The upper end of the instrument is closed with a ground-glass stopper.

When arranged with glass-wool as described, the whole is dried at 110° C., allowed to cool, and weighed.

The filter is fixed in a suitable support, and the fluid with the uric acid precipitate is placed upon it. The filtrate is used to wash out the uric acid from the capsule in which it was formed, and this is repeated until no trace of the uric acid is left in the latter, the whole having been placed on the filter. Finally, the latter is washed repeatedly with a little water, and best by means of an aspirator, after which the filter and precipitate together are dried at 100° C. They are then allowed to cool, and small quantities of bisulphide of carbon added in three portions of about 2 or 3 cc., the bisulphide of carbon removed by the addition of æther, and the filter dried at 100° C. until it attains a constant weight. The difference between this weight and that of the filter as previously ascertained expresses the amount of uric acid in the quantity of urine taken. The dried filter containing the uric acid may be conveniently weighed by placing it in the scale upon a little triangular glass support of known weight and hollowed into a suitable angle, in which the thin end of the filter rests. The disturbing oscillation of the latter in the scale-pan may be prevented in this way.

*Hopkins' Method.*⁵⁶³—This method is preferable to that of *Ludwig* and *Salkowski*. To 100 cc. of urine, powdered ammonium chloride is added to saturation. For this purpose 30 grms. [50 grms. (*Hopkins*)] will suffice. The mixture should be stirred occasionally and allowed to stand in a cool place for twenty-four hours. It is then filtered through a thin filter paper, and the precipitate thereon is washed three or four times with a saturated solution of ammonium chloride, the filter expressed, and the precipitate rinsed off with a little boiling water into a porcelain saucer, treated with 5 cc. of dilute (one-fourth) hydrochloric acid, evaporated to one-half its bulk on the water-bath, allowed to cool, passed through a Ludwig's filter of known weight, and the process described on p. 361 gone through. The increase of weight is that of the uric acid in 100 cc. of urine. The procedure differs from that of *Hopkins* only in substituting an estimation by weight for that of titration with potassium permanganate. Weighing gives more accurate results. Two estimations should be made and the mean taken. This method, besides being simpler than that of *Ludwig* and *Salkowski*, avoids the ambiguity occasionally involved in the latter by the formation of silver sulphide. Other authors, e.g., *Wartapetoso*,⁵⁶⁴ have also come to approximately the same opinion with regard to the value of this method.

A healthy adult excretes 0.2–1.0 grm. of uric acid with the urine in twenty-four hours, or, according to *Herter* and *Smith*,⁵⁶⁵ 0.5–0.75 grm. The quantity is increased in health by an abundant animal diet, and pathologically in fever, leukæmia (*Fleischer* and *Penzoldt*, *Bohland* and *Schurz*⁵⁶⁶), pernicious anaemia, and in diseases of the heart and lungs with obstructed respiration.⁵⁶⁷ A series of observations which the author has made with the *Salkowski-Ludwig* method in a case of diabetes gave the quantity of uric acid excreted as between 0.9400 and 1.4814 grm., and this quantity was not diminished by the administration of alkalies; and a marked increase was noted, while haemorrhage lasted, in a case of scurvy.⁵⁶⁸

The excretion of uric acid is diminished in a number of chronic affections, such as nephritis, gout (after the acute paroxysm), diabetes, and chronic arthritis. A diminution was also found by *v. Bamberger*⁵⁶⁹ in a case of progressive muscular atrophy. *Salkowski* and *Spilker*⁵⁷⁰ have observed that taking alkali internally is followed by a fall in the amount of uric acid in the urine; and in sick children the use of alcohol has a similar effect (*v. Jaksch*).⁵⁷¹ An increase has been noted in feeding with thymus gland.⁵⁷² The observations of *Kuhnau*⁵⁷³ and others have shown that the white blood corpuscles are instrumental in the elaboration of uric acid.

Finally, we sometimes meet with a condition in which, with emaciation and certain subjective symptoms, as hypochondriasis, &c., is asso-

ciated an enormous increase in the elimination of uric acid as the only objective manifestation, and such cases undoubtedly constitute what is called the uric acid diathesis.⁵⁷⁴

XVIII. Urea and Total Nitrogen.—By far the greater part—about 90 per cent.—of the nitrogen taken in with the food is eliminated as urea. It must not, however, be forgotten that there are in the urine other nitrogenous compounds, such as uric and hippuric acids, and other amido-acids, and ammonia salts, but the great bulk of nitrogenous waste is removed in the form of urea. Of this body 32-40 grms. are daily excreted by a healthy man, but its quantity varies considerably under physiological, and still more under morbid conditions.

Amongst diseases, fever and diabetes are attended with increased elimination of urea. [Prout⁵⁷⁵ has described a morbid condition which he calls *azoturia*, and which he ascribes to an excessive formation of urea.] On the other hand, this is diminished in chronic affections accompanied by malnutrition and in diseases involving the proper structure of the liver, where it is elaborated (*Schröder*).

The quantity of urea excreted is lessened by taking alcohol in the case of children (*v. Jakisch*).⁵⁷⁶ On the other hand, there is an increased excretion of urea in children during the febrile period of lobar pneumonia (*v. Jakisch*).⁵⁷⁷ Bernabei⁵⁷⁸ has observed a diminished excretion (*hypozoturia*) as a constant occurrence in chronic alcoholism. Numerous recent observations show that the exhibition of thyroidal preparations frequently results in an increased excretion of nitrogen (*Darid*).⁵⁷⁹

Variations in the output of urea are an expression of changes in nitrogenous metabolism generally, and as such possess the highest clinical interest. It would be well if we could measure the quantity of that body in the urine at any given time, but the processes by which this can be done are not available for clinical purposes. The best and most satisfactory procedure is to estimate the total quantity of nitrogen by *Kjellahl's* method.

An approximate estimate of the quantity of urea excreted in the course of twenty-four hours may, however, be effected by *Hufner's*⁵⁸⁰ method. In this the urea is decomposed by means of alkaline hypobromites, and the nitrogen given off as a gas is collected, whilst the carbonic acid combines with caustic soda present. The apparatus required is represented in the accompanying figure. It consists of a cylinder of stout glass of 100 cc. contents (B), expanding at its middle, and connected below by means of a binder and tight-fitting tap with a smaller tube (A), also of glass, which holds about 5 cc. It is important that the capacity of the latter, which serves to receive the urine, together with that of the perforation in the tap, shall be accurately known. To this end the apparatus is carefully washed with water,

rinsed out with alcohol, and dried. The tap is opened, and mercury poured into the lower vessel so as to overflow into the upper one. The tap is now closed, the overflow of mercury above it poured off, and the contents of A removed and weighed in a vessel whose own weight is known. The result divided by the sp. gr. of mercury (13.59) gives the

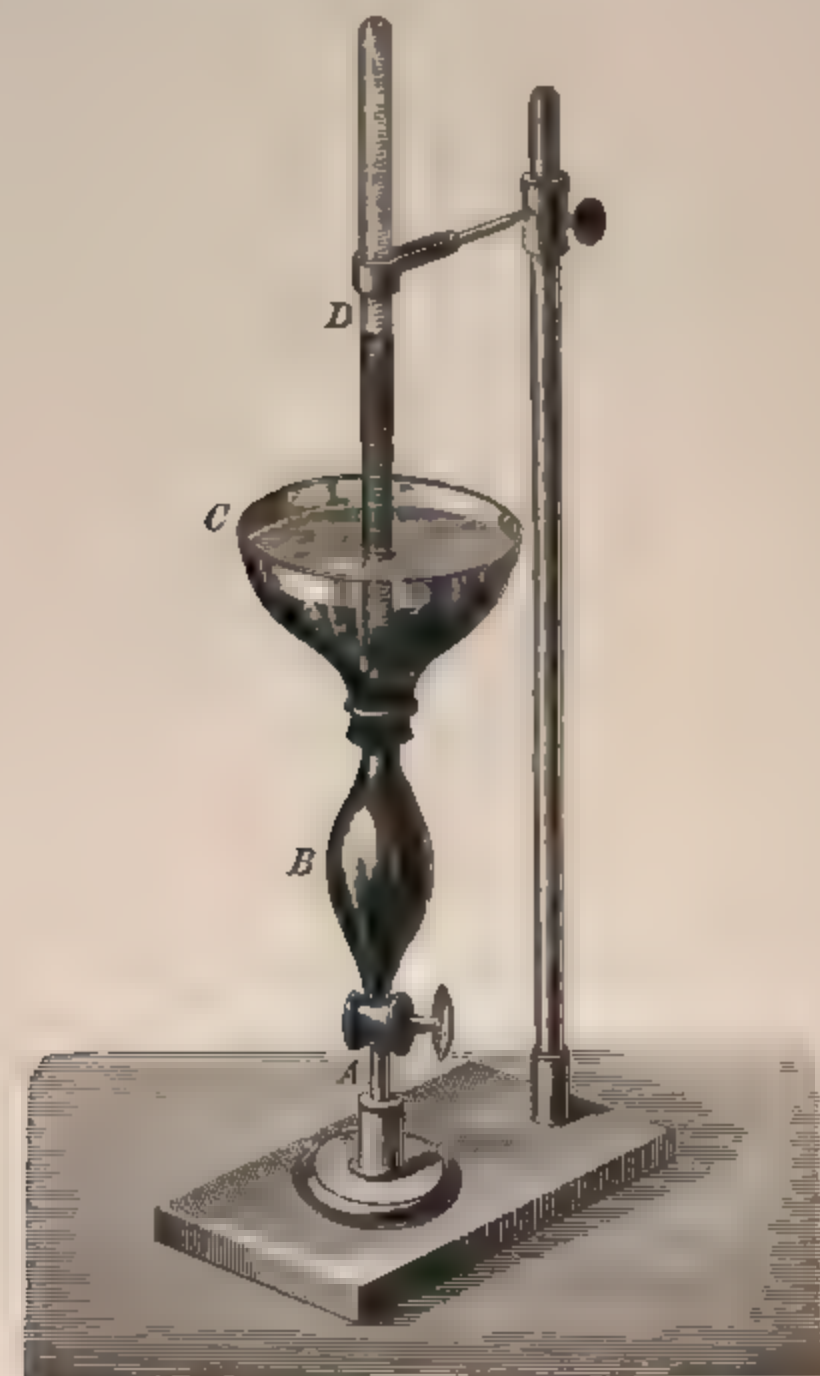


FIG. 140. Hünfer's Apparatus for Estimation of Urea.

cubic contents of the tube A. This must be verified by repeating the process, and if the capacity ascertained should vary each time, the mean must be taken. It should be calculated to three places of decimals.

In cases where the method of weighing cannot be conveniently carried out, the capacity of the vessel destined to hold the urine may be ascertained with great accuracy thus: This portion of the apparatus is filled with a watery solu-

tion of some aniline dye which is not taken up by chloroform, the apparatus washed out with chloroform, and the tinted water together with the chloroform, in which little or none of the dye dissolves, is placed in an accurately graduated burette. The chloroform is allowed to settle and the quantity of the aniline fluid in the burette is read off. This process should be gone through at least three times, and the mean is taken of the ascertained results, which, however, usually agree well.

The remaining parts of the apparatus are a glass bowl (c) fitting by a caoutchouc stopper in its bottom upon the upper extremity of b, and a glass tube 30–40 cm. in length, 2 cm. wide, and accurately graduated in 0.2 cm. units of capacity.

A rough estimate of the proportion of urea is formed either by previous analysis, or, better, by inference from the sp. gr. of the urine, and the latter diluted in such a way that a specimen shall contain not more than 1 per cent. urea. The vessel a, whose capacity is accurately known, is filled by means of a long funnel with urine, the well-greased tap closed, and the bulbous vessel b washed well with water, so as to remove all trace of urine from its surface. Another tube, about half a metre long, may be interposed between the upper end of b and the bowl (c), with the object of prolonging the contact between the column of urine and the hypobromite, and so securing its complete decomposition.

A fresh solution of hypobromite is then made in the following manner:—100 grms. of caustic soda are dissolved in 250 cc. of water, the mixture allowed to cool, and 25 cc. of bromine added. The solution must be freshly prepared for use, withheld from the light, and kept in a cool place. The concentrated fluid so prepared gives better results than the more diluted reagent formerly in use (*Pfluger and Schenck*).⁴⁶¹ With the vessel c in position, b is entirely filled with the above solution, and a concentrated solution of common salt is poured into c to a depth of 1 cm. The graduated tube d is likewise filled with the common salt solution, care being taken to exclude air-bubbles, and inverted in c over the tapering extremity of b, which projects into the solution of common salt contained in that vessel. It is fixed by a clamp in this position. Distilled water may be substituted for the salt solution.

The tap is now opened. The relatively heavier hypobromite solution sinks, and as it does so a rapid evolution of gas takes place and lasts for 15–20 minutes. The nitrogen formed collects in the graduated tube, and when it has ceased to be given off, the latter is closed with the thumb and, still inverted, placed in a cylinder of pure water. It is there held with a clamp, and is depressed as much as possible for a space of fifteen minutes. After that it is raised with wooden forceps until the fluid in the tube and that in the cylinder stand at the same level. The volume of contained gas is then read off, and the barometric pressure and temperature are noted at the same time.

From the volume of nitrogen obtained in this manner the weight in grammes of the urea taken may be deduced by the following formula :—

$$G = \frac{v(b - b')}{354.3 \cdot 760 (1 + c \cdot 00366t)}$$

Where

G = weight of urea in grammes.

v = volume of gas generated (in cc.).

t = temperature.

b = barometric record.

b' = tension of water-vapour at temperature t .

The percentage of urea is expressed by the product of $G \times 100$ divided by the volume of urine analysed. The number 354.3 is substituted in the equation for 372.7, since it has been found that from 1 grm. of urea the total quantity of nitrogen, namely, 372.7 cc., is never obtained in this way, and that the number chosen more aptly represents the fact.

The value of b' will be found in Bunsen's tables, from which the following figures are extracted.⁵⁸² They express in millimetres the tension of water-vapour at the corresponding temperatures, which are those most commonly existent :—

10° C.	9.165	14° C.	11.908	18° C.	15.357	22° C.	19.659
11° C.	9.792	15° C.	12.699	19° C.	16.346	23° C.	20.888
12° C.	10.457	16° C.	13.536	20° C.	17.391	24° C.	22.184
13° C.	11.162	17° C.	14.421	21° C.	18.495	25° C.	23.550

To carry on such investigations uninterruptedly, at least two sets of apparatus should be available. The researches of *Pflüger*⁵⁸³ and his pupils have shown that the results obtained are not entirely accurate, but sufficiently approximate. The method has the advantage over others presently to be described, that it can be carried out quickly. Moreover, the object of a clinical investigation is less often to ascertain the precise quantity of urea than to determine its variations at different times, and this purpose the method admirably fulfils. *Huppert*⁵⁸⁴ has shown that by this process one can ascertain approximately the total amount of nitrogen excreted in the urine, provided that the uncorrected number representing the amount of nitrogen, as obtained by Hüfner's method, be multiplied by 1.136. In the case of a febrile urine the factor is 1.18. Recently a number of other apparatus of the same kind have been employed.⁵⁸⁵ That of *Lange* seems specially serviceable.⁵⁸⁶

The quantitative estimation of urea may also be effected by Liebig's titration method, as modified by Pflüger. It will be found described at length in the systematic works of *Huppert*, *Hoppe-Seyler*, and *Leube-*

*Salkowski.*⁵⁸⁷ The method of *Mörner* and *Sjöqvist*⁵⁸⁸ is also a good one, but the figures derived by it are a little too high.

To estimate exactly the total quantity of nitrogen obtainable from the urinary products, other methods are needed, as those of *Will-Varrentrapp* and *J. Kjeldahl*.⁵⁸⁹

Kjeldahl's Method.—Five cc. of urine are placed in a Kjeldahl flask, a little yellow oxide of mercury is added, and then 10 cc. of concentrated sulphuric acid. The mixture is heated over a flame until all its colour has disappeared. It is then allowed to cool, poured carefully into a flask of about a litre capacity; 40 cc. of a solution of potassium sulphide (40 grms. to the litre of water) are first added, and afterwards 80 cc. of a solution of nitrogen-free sodium hydrate (270 grms. to the litre). The flask is then stoppered.

Gunning's mixture may be employed with advantage. Oxidation occurs then more rapidly, and the addition of potassium sulphide is unnecessary. It has the following composition: 10 grms. pure potassium sulphate, 0.5 grm. cupric sulphate, and 15 cc. pure sulphuric acid. Powdered talc may be added to diminish the force of decomposition.

Two or three pieces of metallic zinc are introduced, the fluid boiled without shaking, and distillation effected. The ammonia driven off is led into a known quantity of $\frac{1}{4}$ -normal sulphuric acid, and the sulphuric acid which remains coloured with *May's* litmus tincture and titrated with $\frac{1}{4}$ -normal soda solution. The number of cc. of $\frac{1}{4}$ -normal acid solution required to neutralise the ammonia produced, multiplied by 0.0035, gives the quantity of nitrogen in 5 cc., and this multiplied by 20 gives the percentage of nitrogen in the urine.

The apparatus required is shown in fig. 141; A leads on and B leads off the current of cold water. The $\frac{1}{4}$ -normal sulphuric acid solution (of which 30 cc. generally suffices) is supplied to the receiver flask in porcelain beads. At the end of distillation, that is, when about two-thirds of the fluid in the distillation flask has been driven off, and the boiling mixture begins to effervesce, the beads and retort tube are rinsed out with water, and titration performed. Two estimations should be made.

It is often desirable to determine the amount of urea with reference to the total excretion of nitrogen, and this may be done by the application separately of *Mörner's* and *Kjeldahl's* methods.

The qualitative tests for urea are described in the chapter on Blood. In connection with urine they possess but little practical interest.

XIX. Kreatinin.—The formation of kreatinin has been shown to be intimately associated with the decomposition of muscle-substance, and the quantity produced is in direct relation to the amount of flesh-meat

consumed as food, and, under certain circumstances, to the muscle-waste within the body. Under these circumstances muscle contains kreatin, which is changed into kreatinin as it passes through the body.

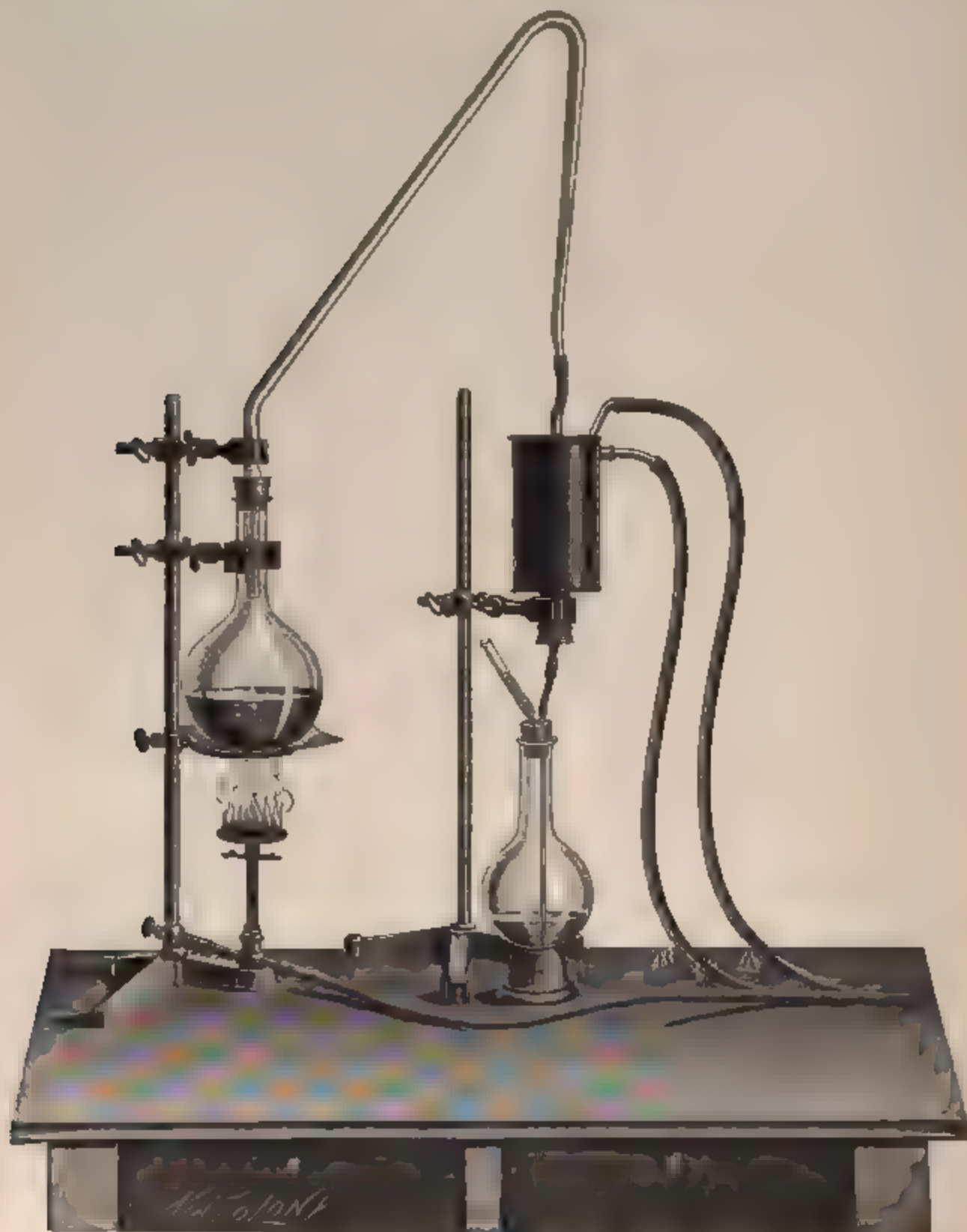


FIG. 141.—Apparatus used for the Distillation of Ammonia in the Kjeldahl Process.

This is to be borne in mind when a clinical inference is drawn from the observation of an increase or diminution of the quantity of kreatinin secreted. Hitherto such inferences as were possible have had but a

limited application to the purposes of diagnosis. They are insufficiently based, and rest for the most part on the experience of individual cases. According to *Neubauer*,⁵⁹⁰ the quantity excreted by a healthy man is about 1 grm. According to *Pouchet*,⁵⁹¹ it is 1 grm. in the case of the male, 0.75 grm. in that of the female, while kreatinin is altogether absent from the urine of sucklings. This point, however, is contested by *Grocco*.⁵⁹²

An increase in the output of kreatinin has been observed in acute diseases of all kinds while attended with fever, and in diabetes (*Senator*),⁵⁹³ and a diminution in chronic nephritis and diabetes insipidus, in convalescence after acute diseases, in chlorosis, anaemia, tuberculosis, and marasmus,⁵⁹⁴ and as a consequence of insufficient feeding.⁵⁹⁵

Kreatinin is a base which forms highly characteristic compounds with acids, such as phosphotungstic and phosphomolybdic, hydrochloric and sulphuric, and with salts of the heavy metals.

Qualitative Tests.—Kreatinin may be detected directly in the urine by the methods of *Weyl*⁵⁹⁶ and *Jaffé*.⁵⁹⁷

Weyl's Test.—For the application of this test the urine should be freed from acetone by distillation (p. 354). A freshly prepared and very dilute solution of nitroprusside of soda and caustic potash is added. If kreatinin be present, the fluid will take a beautiful red colour (like that produced in *Legal's* test for acetone), which soon disappears, and does not return with the further addition of acetic acid.

Jaffé's Test.—A fairly concentrated solution of picric acid and a little caustic potash are added to the urine. If the fluid be heated, the presence of kreatinin will be shown by the appearance of a beautiful red coloration. Acetone and grape-sugar yield a similar reaction. Picric acid with caustic potash alone gives a slight red colour.

Quantitative Estimation.—Kreatinin forms with zinc chloride a double salt of slight solubility, and this property is made the basis of the quantitative method, which was first devised by *Neubauer*⁵⁹⁸ and modified by *Salkowski*.⁵⁹⁹

Two hundred cc. of urine are mixed with a little milk of lime until the fluid has an alkaline reaction. This is to precipitate phosphoric acid. A solution of calcium chloride is added until a precipitate ceases to form. The fluid is allowed to stand for half-an-hour, after which the precipitate is filtered off and repeatedly washed with water, and the filtrate and washings (acidulated with a little sulphuric acid) are evaporated on the water-bath to a syrupy consistence. To the residue is added 50–100 cc. of 78 per cent. alcohol; the mixture is well stirred and allowed to stand in the cold for several (6–8) hours; then it is filtered, and to the filtrate (which, if alkaline, must be rendered acid with acetic acid) 10–15 drops of an alcoholic solution of zinc chloride

are added. The latter is prepared by adding alcohol to a concentrated solution of zinc chloride until a density of 1.2 is attained. After the lapse of two or three days the precipitate is brought upon a filter of known weight, and the filtrate as it passes through is constantly returned to wash the vessel which contained the precipitate. When all the precipitate has been brought upon the filter in this way, it is washed with 90 per cent. alcohol until the filtrate shows but slight opalescence with silver nitrate; after which it is dried to a constant weight at 100° C. One grm. kreatinin-zinc-chloride corresponds to 0.6242 grm. kreatinin. The amount of kreatinin in the quantity of urine taken may therefore be known by multiplying the quantity of kreatinin-zinc-chloride obtained by 0.6242.⁶⁰⁰

*Kolisch*⁶⁰¹ suggests that kreatinin may be precipitated by corrosive sublimate from the urine, and the total nitrogen of the precipitate estimated by *Kjeldahl's* process; from this the quantity of kreatinin may be deduced.

Kreatin, which is closely related to kreatinin, does not occur as such in the urine, but it is readily formed from kreatinin in alkaline fluids. Hence it follows that alkaline urine should not be employed clinically for the quantitative estimation of kreatinin.

XX. Xanthin Substances.—In addition to the substances mentioned in this chapter, *sub. XVIII.*, human urine contains certain other nitrogenous organic compounds, by means of which nitrogen is also eliminated from the system. Amongst these are betain, hypoxanthin (sarkin), xanthin, and xanthokreatinin. The occurrence of these substances is without great clinical significance at present. Certain basic substances have recently been isolated from urine by precipitation with phosphoric acid (*Thudichum*⁶⁰²), such as urochrome, urotheobromin, omichol, and reducin. The physiological action of these substances is not yet known.

*Salomon*⁶⁰³ has proved that hypoxanthin is a normal constituent of urine, and so also are carnin, guanin, paraxanthin, and heteroxanthin. The relations recently shown to exist between cell nuclei and the nuclein bases (xanthin, guanin, hypoxanthin, adenin) [*Kossel*] confer a clinical significance upon the presence in the urine of substances of the xanthin group. This quantitative estimation is effected best by the process of *Krüger* and *Wulff*.⁶⁰⁴ In this process the uric acid and xanthin substances are precipitated together from one portion of the urine by means of cupric sulphate and sodium bisulphide, and the total nitrogen estimated by *Kjeldahl's* method; while from another portion the uric acid is estimated, its nitrogenous constituent computed, and this deducted from the result in the first case; the difference shows the amount of

nitrogen derived from the xanthin bases, and from this the quantity of the latter in a given volume of urine is deduced. The details are as follows :—100 cc. of urine are taken, and the uric acid is determined by Hopkins' method. Two estimations should be made, and the mean result divided by three. This gives the quantity of nitrogen combined as uric acid. Another 100 cc. freed from albumin is heated to boiling, and to it is added 10 cc. of a 50 per cent. sodium bisulphide solution and 10 cc. of a 13 per cent. cupric sulphate solution, when the mixture is again boiled. Five cc. of a 10 per cent. barium chloride solution are next added. The resulting precipitate is allowed to stand for two hours, filtered off, washed with boiled water at 60° C., and submitted to *Kjeldahl's* process (see p. 367). *Krüger* and *Wulff* use Gunning's mixture (see p. 367). The quantity of nitrogen derived less than ascertained as the component of uric acid, belongs to the xanthin substances, and this multiplied by

$\frac{100}{36.295}$ or 2.755 gives the absolute amount of the latter in 100 cc. of

urine. A mixture in equal quantities of xanthin, guanin, hypoxanthin, paraxanthin, heteroxanthin, and carnin, contains 36.295 per cent. of nitrogen. Further experience is needed to show the relation between the elimination of these bodies and definite diseased states.⁶⁰⁵ Already many interesting particulars have been recorded.⁶⁰⁶

It may be mentioned that nitrogen is set free in the system under certain circumstances, as after the ingestion of calcium hydrate in the form of carbamates (*Abel* and *Muirhead*).⁶⁰⁷ According to *Ludwig* and *Savoy* puerperal eclampsia is probably due to the formation of carbamic acid within the body.

XXI. Ptomaines (Putrefaction Bases) in the Urine.—It would appear from the investigations of *Pouchet*⁶⁰⁸ that healthy urine contains traces of certain toxic substances of an alkaloid character, and according to the researches of *Bouchard*,⁶⁰⁹ *Lépine*, and *Guerin*,⁶¹⁰ these bodies are more abundant under morbid conditions. They were found by *A. Villiers*⁶¹¹ as an invariable manifestation in measles, diphtheria, and pneumonia; and in the urine of cholera *A. G. Pouchet*⁶¹² discovered an alkaloid which was not identical with that observed by him in the faeces of the same disease (see p. 243). *Feltz*⁶¹³ found similar bodies in the urine of cancer patients, and *Lépine*⁶¹⁴ in that of pneumonia. *Roges* and *Gauze*⁶¹⁵ observed that the toxic property of the urine was lessened in the febrile period of pneumonia (retention of potash salts?). Observations of this kind have lately been very numerous. Toxines have been found in the urine of scarlatina and pneumonia (*Albu*); in gastric cancer and Addison's disease (*Eicald*, *Jacobsen*);⁶¹⁶ in pleurisy, influenza, and cancer (*Griffiths*).⁶¹⁷ *Bouchard*⁶¹⁸ discovered that human urine acted as a poison when injected within the veins of animals

(rabbits), and he referred the effects to various substances, among which were animal alkaloids. For the detection of the urinary alkaloids the following method has been adopted by *Tanret*, *Bouchardat*, and *Cardier*:⁶¹⁹—

To the urine which has been acidulated with acetic acid a solution of the iodide of mercury and potassium is added. The precipitate, which contains the alkaloids, is readily distinguished from those of other substances, as albumin, mucin, and uric acid, obtained with the same reagent, by its solubility in alcohol at a warm heat.

Ch. Bouchard rendered the urine alkaline with caustic soda, and derived a poisonous body as an æthereal extract.

Pouchet fixed the alkaloid by combination with tannic acid, and subsequently precipitated it with oxide of lead from alcoholic solution.

The methods adopted by the other observers mentioned above differed much as to details. They will be found described at length in the original communications. That of *Brieger* (p. 187) serves best for the detection of animal alkaloids in the urine. In some cases, however, it is important that the urine be previously concentrated in vacuo. Should no result be obtained with this, *Gautier's*⁶²⁰ method may next be tried. Finally, the Stas-Otto method may be applied to the same purpose. (See p. 187.)

The diamines of the urine may be precipitated as benzoyl compounds, and best by the action of benzoyl chloride and caustic potash. *Baumann* and *Udransky*⁶²¹ succeeded in separating several basic derivatives, amongst them cadaverin (pentamethylendiamine), putrescin (tetramethyl diamine), and a small quantity of a third diamine, from the urine of a patient with cystinuria and vesical catarrh. Normal urine was found to be free from these bodies. The author has been for some time engaged in observing the occurrence of similar poisonous bases in the urine of health and disease, and he has found that normal urine and that of some diseases, notably typhoid, pneumonia, leukæmia, cystic pancreas, and Weil's disease,⁶²² hold such only in very small quantity. He would venture to make some suggestions for the benefit of those who are engaged with similar researches. In the first place, it would be well to follow the example of *Brieger*, *Baumann*, and *v. Udransky*, and withhold the name of alkaloids from the bodies (diamines) alluded to, which are derived from the system under morbid conditions, because all that have been recognised as yet are simply diamines, and because none yet examined exhibit the characteristic property of alkaloids, namely, the pyridin radicle. Next, it would be desirable to discriminate between the physiological bases of the urine (kreatinin, reducin, &c.), which belong normally to the fluid, and those which are associated only with certain diseased states. It is not intended to imply that the physiological bases

cannot under any circumstances give rise to the symptoms of disease or of poisoning. (See below.) Experience is not wanting to make it seem in the highest degree probable that the retention, and still more the increased formation, of such physiological products in certain diseases may induce symptoms of the gravest character, and greatly imperil the life of the patient.

Again, it would appear that in certain acute affections specific substances of a toxic character, not observable in normal urine, may be excreted with that fluid. Undoubtedly the matter is somewhat obscure. The author's views may be stated as follows.⁶²³ It is possible to distinguish:—

(1.) Clinical (morbid) symptoms depending upon the retention of the physiological bases (and under this heading would come uræmia) and certain of the symptoms of obstruction (*retention toxicosis*).

(2.) Clinical symptoms, referable to the presence of basic products, which are formed in the system (blood, &c.) in disease and eliminated with the urine (*auto-toxicosis*). [Bruschettini⁶²⁴ has conveyed tetanus to animals (rats and rabbits) by injecting the urine of patients suffering from that disease a fact which would make it appear that the poison of tetanus is eliminated by the urine.]

(3.) Clinical symptoms which are caused by the formation of toxic basic substances from morbid matter, such as pathological fluids lodged in certain parts of the system. Such bases are absorbed, and then give rise to manifestations of severe poisoning. Under this heading would come the collective symptoms of ammoniæmia (see p. 93), and others which follow the absorption of gangrenous pus (*auto-toxicosis*). The latter are sometimes characterised by the presence of guanin,⁶²⁵ and it is probable that the formation of toxalbumins has much to do with them. These may readily be detected by the methods of Brieger and Fraenkel.⁶²⁶

(4.) Clinical symptoms, and consequently morbid types, induced by the action of toxic bases taken into the system with the food, such as the poison of sausages and cheese, p. 189 (*exogenous toxicosis*).

These distinctions are based partly upon clinical observation and partly upon experiments on animals. They will serve as a scheme for the further elucidation of this very important subject.

If we have dwelt at some length on the result of observations which are not yet completed, we have done so because we believe that the careful investigation of the urine in this direction will throw light upon the nature of some diseases which are at present not sufficiently understood.⁶²⁷

XXII. The Ferments of the Urine.—The appearance in the urine of a body resembling pepsin was long ago established by v.

*Brücke.*⁶²⁸ *Sahli, Leo, Gehrig, Stadelmann, and Patella*⁶²⁹ made similar experiments, and confirmed the presence of pepsin in the urine. It has been asserted that trypsin is also an occasional constituent of that fluid, though some observers, *Leo, Stadelmann, and Grützner*,⁶³⁰ have failed to find it.

With reference to the pepsin ferment of the urine, clinical interest attaches to it from the fact that it is absent in typhoid and carcinoma of the stomach,⁶³¹ and according to some observers (*Mya, Belfanti*⁶³²) in nephritis.

For the detection of pepsin *Sahli's* method (adapted from those of v. Wittich and Grützner) may be employed. It is founded upon the property which, as v. Wittich originally observed, blood fibrin possesses of readily absorbing that body from solutions. To this end a little pure fibrin is placed in the urine and allowed to rest there several hours. It is then removed, placed in dilute hydrochloric acid (.2 per cent.), and the mixture kept at a temperature of 30-40° C. Any pepsin present is precipitated on the fibrin, and the latter is slowly digested in the acid fluid.

Diastatic ferment also is said by *Hovoltschiner* and *Rosenberg* to occur in urine.⁶³³ The researches of *Breusing*⁶³⁴ and observations made by the author tend to show that the ferment in question is in many cases not diastase, but an amylolytic substance. At the same time it must be stated that the author has frequently determined the presence of diastase by the usual methods, both in health and disease. So, too, *Leo*⁶³⁵ has found diastase both in healthy and in morbid urine.

Milk-curdling ferment is occasionally present in the urine (*Hovoltschiner*,⁶³⁶ *Boas*,⁶³⁷ *Helvæs*).

As to whether the urine contains a ferment capable of decomposing urea into carbonic acid and ammonia, opinions differ. On the one hand, *Musculus*⁶³⁸ believes that he has isolated such a ferment, but *Leube*⁶³⁹ has sought for it in vain in urine which was actually undergoing ammoniacal fermentation.⁶⁴⁰

B. Inorganic Substances.—The inorganic constituents of the urine are for the most part salts of hydrochloric, sulphuric, and phosphoric acids, to which must be added carbonates, silicates, nitrates and nitrites, and sulphuretted hydrogen.

1. Chlorides.—The chlorides of sodium, potassium, ammonium, and magnesium⁶⁴¹ are found in the urine, and of these we are chiefly concerned with chloride of sodium. Of this salt 10-15 grms. are voided by a healthy man in twenty-four hours; but its quantity is greatly influenced, even in disease, by the supply of salt taken as food.

It is increased by an abundant diet and as a consequence of conditions which determine the retention of chlorides within the system;

and diminished in fevers, and notably in croupous pneumonia.⁶⁴² The elimination of chloride of sodium has also been observed to be less in cases of chronic nephritis, and sometimes in certain diseases of the stomach (*Gluzinski*).⁶⁴³

According to *Huchard*,⁶⁴⁴ the quantity of chlorides falls as low as 2-3 grms. in acute diseases, as well as in cases of pneumonia and diarrhoea. A reduction to 2 grms.—independent of the quantity of nutriment absorbed—is an unfavourable indication in chronic diseases; in fact, the cessation of chloride excretion in such cases foreshadows the approaching death of the patient.

Detection of Chlorides.—The urine is treated with nitric acid, and a solution of nitrate of silver added. A caseous precipitate soluble in ammonia, insoluble in nitric acid, shows the presence of chlorides.

Quantitative Estimation of Chlorides.—*Mohr's method* is to treat the urine with chromate of potash, and gradually add nitrate of silver, when all the chlorine combines with the silver to form silver chloride, and the occurrence of a red precipitate (chromate of silver) marks the end of the reaction. The details of the process will be found in works on urinary chemistry. *Salkowsky's*⁶⁴⁵ modification of *Volhard's*⁶⁴⁶ method is to be preferred.

When to a solution of nitrate of silver acidulated with nitric acid is added some of a solution of sulpho-cyanide of ammonium, a curdy white precipitate forms, and this, like chloride of silver, is insoluble in nitric acid, soluble in ammonia. If the fluid also contains a ferric salt, a blood-red colour (ferrocyanide) forms at the moment when the last of the silver is precipitated. If now the sulpho-cyanide of ammonium solution is of a known degree of concentration, it is possible to determine the quantity of silver present by noting the point at which the red coloration takes place. To apply this principle for gauging chlorides in solution to the fluid containing them, a silver solution of a definite degree of concentration is added in excess, and that portion of it which is not precipitated as chloride of silver is measured in the manner indicated above.

The following solutions are needed in the process:—

- i. *Pure nitric acid of 1.2 sp. gr.*
- ii. *Concentrated solution of double sulphate of iron and ammonia free from chlorine.* It is necessary that this be free from chlorine, and if not already so, it must be purified by crystallisation.
- iii. *Nitrate of silver solution of definite strength.* The chemically pure crystalline salt is dissolved in water in the proportion of 29.075 grms. to the litre. A cc. of this solution corresponds to 0.01 grm. of chloride of sodium.
- iv. *Sulpho-cyanide of ammonium solution.* This should be of such

a strength⁶⁴⁷ that 25 cc. shall correspond to 10 cc. of the silver solution. For this purpose 6.5 - 7 grms. of sulpho-cyanide of ammonium may be dissolved in water, and more water added to 400 cc. A burette is filled with the solution so prepared.

Titration with the sulpho-cyanide of ammonium solution is effected thus:—Ten cc. of the silver solution (iii.) is placed in a flask and diluted with water to 100 cc.; 4 cc. of nitric acid (i.) are next added, and after that 5 cc. of the double sulphate of iron and ammonia (ii.). The mixture is well shaken up, and sulpho-cyanide of ammonium solution from the burette is carefully added until a slight but permanent red coloration appears. The process is repeated several times, the quantity of the reagent employed in each case being noted and the mean taken.

In accordance with the result obtained, the sulpho-cyanide of ammonium solution is diluted to such a point that 25 cc. shall correspond to 10 cc. of the silver solution. Thus, if the terminal reaction (red colour) occurs after the addition of 22 cc., the following formula may be applied to determine the volume to which a litre of the solution must be diluted:— $22 : 25 = 1000 : x$, and $x = 1136.3$. To the litre of sulpho-cyanide of ammonium, therefore, 136.3 cc. of water must be added in order that 25 cc. shall correspond to 10 cc. of the silver solution (iii.).

The remainder of the process is as follows:—Ten cc. of urine are measured out with a pipette and placed in a graduated flask of 100 cc. capacity; 50 cc. of water are added, and then successively 4 cc. of the nitric acid (i.) and 15 cc. of the silver solution (iii.). The flask is closed with a glass stopper, and well shaken up until a precipitate ceases to form and the fluid tends to clear; the flask is then filled to the mark 100; its contents passed through a dry paper filter and received into a dried measure cylinder or a flask holding 80 cc. The 80 cc. of fluid thus obtained are poured into a larger flask of some 250 cc. capacity; 5 cc. of the double sulphate of iron and ammonia (ii.) are added, and the sulpho-cyanide ammonium solution (iv.), prepared in the manner already indicated, is gradually supplied from a burette, until the terminal reaction is shown on shaking the fluid by a faint but abiding red coloration. The quantity of the sulpho-cyanide of ammonium used to effect this is now read off. Experience has shown that 15 cc. of the silver solution is more than sufficient to precipitate all the chlorine from urine strongly acidulated with nitric acid, and that an excess of silver nitrate remains in solution. The excess of silver may be measured volumetrically by means of the sulpho-cyanide of ammonium solution, and the quantity of chlorine in the urine calculated from the difference.

The quantity of chloride of sodium in grammes in one litre of the urine may be ascertained by the following formula :—

$$x = [37.5 - \frac{5}{4} R] \frac{4}{10}$$

Where

x = the quantity of NaCl in a litre of urine in grms.

R = the quantity of sulpho-cyanide solution used in cc.

The formula is derived thus :—

Ten cc. of the silver solution correspond to 25 cc. of the sulpho-cyanide of ammonium, and consequently 15 cc. of the former to 37.5 cc. of the latter. For 100 cc. of the fluid examined, therefore, are needed 37.5 cc. of the sulpho-cyanide solution less by five-fourths of the total quantity of the latter employed, since 80 cc. of the titration fluid, i.e. sulpho-cyanide, was used ; consequently 100 cc., being the volume of the original fluid, will require five-fourths of the quantity read off (on the burette).

Now 25 cc. of the sulpho-cyanide correspond to 10 cc. of the silver solution, and therefore 1 cc. of the former to 0.4 cc. of the latter.

One cc. silver solution corresponds to 0.01 grm. sodium chloride ; 0.4 cc. silver solution corresponds to 0.004 grm. sodium chloride.

Therefore, to obtain the quantity of chloride in the volume (10 cc.) of urine tested, the expression $[37.5 - \frac{5}{4} R]$ must be multiplied by 0.004, or by $0.4 = \frac{4}{10}$ for 1000 cc. (litre) of the urine.

2. Sulphates.—Sulphuric acid is present in the urine both as simple (preformed) sulphuric acid and as æther-sulphuric (compound sulphuric) acid (see p. 343). The combinations of the latter acid have been already spoken of. Further, the urine contains sulphur as sulpho-cyanides,⁶⁴⁸ hypo-sulphites (thio-sulphates⁶⁴⁹), and sulphuretted hydrogen (see also p. 381).

The total quantity of sulphuric acid excreted in twenty-four hours by a healthy adult under ordinary conditions of diet is about 2 grms., of which 0.1 grm. is in the form of æther-sulphuric acid compounds.

Sodium, potassium, magnesium, and calcium sulphates are found in the urine (see pp. 282, 287). But little clinical significance attaches to a general increase or diminution in the output of sulphuric acid in disease. Of far greater import are changes of the relative quantities of simple and of æther-sulphuric acids (see p. 343).

Thus a urine rich in indigo compounds contains little of the preformed sulphuric acid, and in carbolic acid poisoning this may entirely disappear.

Detection of Simple Sulphuric Acid.—The urine is filtered if turbid, acidified with acetic acid, and solution of chloride of barium added. A fine precipitate, barium sulphate, forms. This reaction never fails with the normal fluid.

Estimation of Simple Sulphuric Acid.—This may be effected by determining first the total quantity of sulphuric acid present, and then that of æther-sulphuric acid, according to the methods indicated at p. 347. The difference will be the quantity of free sulphuric acid sought.

Estimation of the Total Quantity of Sulphur.—Where it is of consequence to ascertain the total quantity of sulphur in the urine, the best plan is to evaporate the alkaline urine, either the whole or a known proportion of it, on the water-bath, then to fuse the incinerated residue with saltpetre and soda (*Heffter*⁶⁵⁰), to extract the fused mass with boiling water, and to proceed further as directed in the process for determining the total quantity of sulphuric acid on p. 348. The extract is treated with barium chloride, and the sulphur estimated as barium sulphate.

3. Phosphates.—The phosphoric acid of the urine is combined partly with sodium, potassium, and ammonium, and partly with lime and magnesia. Being a tribasic acid, it forms three classes of salts—acid, neutral, and basic. Of these, the acid phosphates of alkalies and alkaline earths and the neutral and basic phosphates of the alkalies are soluble in the urinary fluid. The neutral phosphates of the alkaline earths are but little soluble therin, and their basic phosphates still less so.

Upon this fact depends the deposition of phosphatic sediment when the urine is boiled. The acid and neutral phosphates of the alkaline earths are changed in the process into the insoluble basic salts. The phosphorus salts occur partly in solution and partly as a crystalline deposit (see pp. 281, 288).

Two to three grms. of phosphoric acid are excreted in twenty-four hours. According to *Lennmalm*,⁶⁵¹ the amount is the same, in proportion to the body weight, in children as in adults.

It appears, especially from the researches of French authors,⁶⁵² that in certain morbid states the phosphatic constituent undergoes notable increase, and that consequently a condition of phosphaturia, analogous to that of oxaluria, may properly be spoken of: and further, that this condition may take the place of glycosuria in diabetes. The subject, however, is still under discussion. A diminished elimination of phosphates was observed by *Stokris*⁶⁵³ in arthritis; and *r. Jaksch*, unlike other observers, has found that in some, though not all cases of lobar pneumonia amongst children, the quantity of phosphoric acid eliminated during the continuance of fever was increased as compared with the non-febrile period.⁶⁵⁴ *Lennmalm*⁶⁵⁵ has studied this subject in connection with children.

A phosphatic sediment does not imply phosphaturia.⁶⁵⁶ The diagnosis of this condition can be safely based only upon the quantitative

estimation of phosphoric acid, and this is effected best by *Neubauer's*⁶⁵⁷ method of titration with a solution of uranium oxide (see below).

Detection of Phosphates.—The urine is treated with caustic potash and heated. The phosphates are precipitated as earthy phosphates. By the addition of ammonia they may be precipitated without heat.

To detect the presence of phosphoric acid in combination with alkalies, the urine is treated with ammonia and filtered, and to the filtrate an ammoniacal solution of magnesia (a mixture of sulphates of magnesia and ammonia) is added, whereby the phosphates are precipitated as triple phosphate.

Another method is to treat the filtrate (*vide supra*) with acetic acid, when the further addition of uranium solution yields a yellowish-white precipitate.

The same filtrate with perchloride of iron solution gives a white precipitate, which becomes yellow on the addition of more of the perchloride.

Estimation of Phosphoric Acid.—To urine which contains the phosphates as acid phosphates, uranium acetate or nitrate is added in solution until an excess of the reagent first becomes appreciable. If the nitrate be used, free nitric acid forms, and causes a part of the precipitated uranium phosphates to redissolve. To prevent this, in practice a little sodium acetate is added to the urine before titration with uranium nitrate. As an indicator a little tincture of cochineal is employed. This yields a green precipitate in presence of a uranium salt in excess. [Instead of the cochineal fluid a solution of potassium ferrocyanide (1 in 10) may be used. This reagent deposits a deep-brown precipitate with a mere trace of a uranium salt.]

This test, however, is less sensitive in presence of acetate of soda than in simple watery solutions. Hence it is necessary to use a definite quantity of the salt, and to take care that the proportion is maintained in preparing the titration fluid.

The solutions required for the process are:—

i. *Solution of Acetate of Soda.*—A hundred grms. of acetate of soda are dissolved in 800 cc. of water, 100 cc. of a 30 per cent. solution of acetic acid added, and the mixture made up to a litre. Five cc. are employed with 50 cc. of urine.

ii. *Cochineal Tincture.*⁶⁵⁸—A cold infusion is made of a few grms. of cochineal in a quarter of a litre of a fluid composed of 3–4 parts of water with 1 of alcohol, and the solution filtered for use.

iii. *Solution of Uranium Oxide.*—About 20.3 grms. of commercial uranium oxide purified and well dried is dissolved in pure acetic acid, or in the smallest possible quantity of nitric acid, and the preparation made up to a litre. Of the mixture 1 cc. indicates 5 mgrms. of P_2O_5 .

iv. *A Solution containing a Definite Quantity of Phosphoric Acid.*—Fifty cc. should contain precisely 0.1 grm. P_2O_5 . The preparation is made by dissolving 10.085 grms. of neutral phosphate of soda in a litre of water. The commercial salt should be crystallised from solution to obtain it free from chlorine, so that no precipitate forms with nitrate of silver and nitric acid. The crystals are then placed on paper in a funnel, the neck of which is stopped with glass wool, and allowed to dry there until the mother liquid is no longer found to adhere to them. A known weight is then taken and rubbed up in a mortar and a portion of the powder submitted to a gentle heat in a platinum crucible, and finally incinerated. 266 grms. of sodium-pyrophosphate ($Na_4P_2O_7$) correspond to 716 grms. $Na_2HPO_4 + 12H_2O$. Consequently that quantity of the dried crystals which, when incinerated, yields 266 grms. corresponds to 716 grms. of pure phosphate of soda.

Titration Process.—Fifty cc. of the phosphoric acid solution (iv.) are measured into a flask, 5 cc. of the solution of acetate of soda (i.) and a few drops of the cochineal tincture are added, the mixture boiled, and the uranium solution (iii.) gradually supplied until the mixture becomes slightly but permanently green on shaking. In the process a high temperature should be maintained, to promote the formation of uranium phosphates. [When ferrocyanide of potassium is used as the indicator, the addition of the uranium solution is suspended when a precipitate ceases to form. The fluid is again heated, and a drop is tested by adding to it a drop of ferrocyanide in a porcelain capsule. The further supply of uranium solution is regulated by the earliest appearance of a brown colour in the specimens successively tested.]

The uranium solution is now diluted, according to the quantity found to be necessary, as above, in such proportion that 20 cc. shall just suffice for the titration of 50 cc. of the phosphoric acid solution.

Now, 50 cc. of the phosphoric acid solution represent 0.1 grm. P_2O_5 , and consequently 20 cc. of the diluted uranium solution also correspond to 0.1 grm. P_2O_5 .

The titration process is repeated with the urine in precisely the same manner as before: 50 cc. are taken, 5 cc. acetate of soda and a little cochineal added, and the mixture heated and the terminal reaction sought.

Every cubic centimetre of the uranium oxide solution employed in titration represents 5 mgrms. P_2O_5 . Hence the phosphoric acid contained in 50 cc. of urine may be calculated by multiplying the number of cubic centimetres of uranium oxide solution used by 0.005. The result is the quantity of phosphoric acid in grammes contained in 50 cc. of urine. It is advisable in each case to make two such investigations and to take the mean of their results.

4. Carbonates.—The carbonates of lime, magnesia, and ammonia are sometimes present in the urine. The latter salt, however, is found in large quantity only as a result of alkaline decomposition. *Heintz* has proved that ammonium salts may be detected in every specimen of urine, whether decomposed or not. They may be shown best by the addition of milk of lime in a test-tube, when ammonia is given off, and if a piece of red litmus paper be moistened and held over the mouth of the test-tube, it becomes blue. Ammonium carbonate in considerable quantity occurs only in decomposing alkaline urine (see p. 256). The method described in Chapter IV. will serve for the quantitative estimation of ammonia, in which *Nencki* and *Zaleski*⁶⁵⁹ have recently made some very judicious modifications.

Test.—The presence of carbonates in the urine is shown by the evolution of a colourless gas on the addition of acid, and this gas will render baryta water turbid.⁶⁶⁰

5. Nitrates and Nitrites.—Nitrates and nitrites occur in the urine (*Schönbein*).⁶⁶¹ Nitric acid is thought to be derived from the water and food ingested (*Röhmann*).⁶⁶²

Nitrites occur in decomposing urine, and are derived from the reduction of nitrates in urinary fermentation. *Richter*⁶⁶³ has discovered these salts in fresh urine of patients suffering from acute gastric and intestinal catarrh.

Tests for Nitrites:—

(a.) Solution of iodide of starch paste in presence of dilute sulphuric acid or zinc and starch iodide, or the methods described at p. 98.⁶⁶⁴

(b.) Metadiamido-benzol is coloured a deep yellow by nitrites.

(c.) *Karplus*⁶⁶⁵ employs the acetic acid and ferrocyanide of potassium test (*Schäffer's* nitrite reaction). This yields a deep-yellow colour.

*Strümpell*⁶⁶⁶ found hyposulphurous acid in a case of typhoid. The urine became turbid from the separation of sulphur on the addition of hydrochloric acid. The plan adopted by *Salkowski* and *Preisch*⁶⁶⁷ is to distil the urine with hydrochloric acid, when a deposit of sulphur takes place in the upper part of the condensing tube. If sulphur be present in small quantity, this has the appearance of a faint blue exhalation. It should be mentioned further that urine contains traces of silicic-acid (*Pfeiffer*,⁶⁶⁸) and iron salts.

6. Sulphuretted Hydrogen (Hydrothionuria).—Sulphuretted hydrogen is rarely present in the urine, but it can always be obtained from it by heating with mineral acids.⁶⁶⁹ It has been ascertained (*Betz, Senator, Ottario Stejano*⁶⁷⁰) that when retained in the system in considerable quantity it may produce toxic effects (auto-intoxication). In the great majority of cases, according to *Müller*⁶⁷¹ and others (*Rosenheim, Gutzman, Karplus*), hydrothionuria is due to a sulphuretted

hydrogen fermentation of the urine, caused by the action of certain micro-organisms. In one instance of hydrothionuria, diplococci were obtained from the urine, which failed to stain by Gram's method, and these had the property of setting up sulphuretted hydrogen fermentation in sterilised normal urine. *Savor*⁶⁷² met with this condition in a case of prolonged eclamptic coma. An instance of hydrothionuria induced by the incursion of *Bacterium coli*, was published by *v. Stransky*⁶⁷³ from the author's clinic.

This gas is sometimes derived from the alimentary canal, and its presence depends upon an abnormal communication between the urinary passages and the gut. *Betz* further maintains that it may pass by endosmosis from the intestine into the urine; and it would appear also that the gas may be absorbed from the intestine by the blood, and so find its way into the urine. This, according to *Müller*, is a rare event, and happens only when the quantity of sulphuretted hydrogen is so great as to give rise to symptoms of general poisoning.

Tests.—The urine, which should be acid, is placed in a flask, which is closed by a tight-fitting cork. From the latter depends a strip of blotting-paper soaked in sugar of lead and caustic soda. If sulphuretted hydrogen be present, the paper turns black.

Fr. Müller recommends the following plan:—A current of air is passed through the urine, and directed by means of a fine-pointed glass tube upon a strip of blotting-paper soaked in alkaline sugar of lead solution. If sulphuretted hydrogen be present, the paper is blackened. *Emil Fischer's*⁶⁷⁴ test is also applicable to the urine. Some particles of p-amido-dimethylanilin are added to a few cc. of water, and a little concentrated sulphuric acid and one or two drops of a yellow solution of perchloride of iron supplied. The reagent is poured on the surface of the urine to be tested, when, if sulphuretted hydrogen be present, a blue ring (methylene-blue) forms at the place of contact of the two fluids. This ring often takes some minutes to develop. According to *Karplus*,⁶⁷⁵ mercaptan is sometimes found in urine.

7. Peroxide of Hydrogen.—This body was first observed in the urine by *Schönbein*.⁶⁷⁶ Its presence there has no pathological import.

Test.—Dilute solution of indigo is bleached by peroxide of hydrogen in presence of sulphate of iron solution.⁶⁷⁷ Tetra-paper (see p. 160) immersed in the fluid will show the presence of ozone by taking a blue colour.

8. Gases of the Urine.—The urine contains a small proportion of gases, which may be withdrawn from it by the air-pump. They are chiefly carbonic acid, oxygen, and nitrogen.⁶⁷⁸ In rare instances gases have been voided in considerable quantity with the urine, and this has been due to a morbid communication between the alimentary canal and

the urinary passages. Decomposition within the bladder is another cause of gas-formation. *Fr. Müller*⁶⁷⁸ records the case of a man of sixty, with glycosuria and cystitis, where the urine held hydrogen, nitrogen, carbonic acid, and probably methan. *Senator*⁶⁷⁹ has published a similar case.

IV. CHARACTERS OF THE URINE IN DISEASE.

I. The Urine in Febrile States.—In fever the urine is diminished in quantity, acid, deeply coloured, and of high sp. gr. On standing, it often deposits an abundant sediment of urates. Microscopically it exhibits a profusion of crystals of uric acid and urates, and a few hyaline casts, with scattered leucocytes, renal epithelium, or fungi on their surface. It commonly contains a small quantity of albumin (febrile albuminuria) or acetone in variable proportion. Diacetic acid may be present when the disease is of an acute infectious character, or when its subject is a child. In the first case it betokens great danger; not so in the latter.

The presence of peptone (p. 306), with or without serum-albumin and in conjunction with the clinical symptoms of a puerperal or hæmatogenous origin, indicates the formation and absorption of pus within the system.

According to *Ehrlich*,⁶⁸⁰ it is characteristic of the urine in typhoid, measles, and acute tuberculosis to yield a deep-red colour with diazo-benzol-sulphonic acid. Authorities differ much as to the diagnostic value of this reaction. On the one hand, the opinion of *Ehrlich* is supported by *E. B. Goldschmidt*,⁶⁸¹ while *Penzoldt*,⁶⁸² *Petri*,⁶⁸³ *Zaniboni* and *Tessari*⁶⁸⁴ dissent from it. *Ehrlich* obtains the reaction, not with diazo-benzol-sulphonic acid itself, but with sulphanilic acid. Fifty cc. hydrochloric acid are made up to 1000 cc. with water, and sulphanilic acid added to saturation. To 200 cc. of the mixture 5 cc. of a $\frac{1}{2}$ per cent. solution of sodium nitrite are added, and the resulting fluid is added to the urine in equal parts. The mixture is then rendered alkaline with ammonia. *Ehrlich*⁶⁸⁵ has recently recommended that five to six times the volume of absolute alcohol should be added to the fluid to be tested, and the reagent, prepared as above, then added drop by drop to the filtrate. Normal urine gives a yellow colour, while the urine of fever patients, &c., turns scarlet.⁶⁸⁶ The author's experience would induce him to *disclaim for this test any clinical importance whatever, and he could especially enjoin the necessity of avoiding inferences based upon the appearance of the reaction indicated.* He believes that the colour when obtained is always due to the presence of acetone, and he prefers to regard the process rather as an uncertain indication of

that body than as a test for anything else.⁶⁸⁷ The observations of *L. Munson* and *Horst Oertel*⁶⁸⁸ prove that the reaction in question depends always upon the presence of diacetic acid.

It will be seen from this brief statement that a careful analysis of the urine will serve to make evident the details of acute processes earlier and more readily than the methods formerly at our disposal. Other lines of investigation are appropriate to certain acute affections; thus, for instance, in pneumonia and malaria⁶⁸⁹ the presence of chlorides will be sought.

II. The Urine in Disorders of the Circulation (Congestion).

—Under such conditions the urine in its physical characters closely resembles that of fever. It is diminished in quantity, of acid reaction, and high sp. gr. (1.025–1.035). It commonly deposits urates.

Chemically it may be distinguished from fever urine by—

- (1.) The absence of acetone and diacetic acid.
- (2.) The presence of albumin in greater quantity.

Microscopically, and especially when the congestion is chronic, the urine exhibits some leucocytes and altered red blood-corpuscles, often also hyaline casts and cylindrical aggregations of urates (see p. 264, fig. 100), waxy and a few granular casts and renal epithelium. Such constituents indicate secondary changes in the kidney of a chronic inflammatory character.

[The Urine of Phthisis.—*Hale-White*⁶⁹⁰ has pointed out that the urine of phthisis exhibits two peculiarities. Such urine remains acid for a very long time, occasionally for several months, and contains few bacilli, but a large proportion of yeast-like organisms. The persistence of acidity is probably due to a form of acid fermentation. The urine, when kept for a long time, turns very dark in colour, some specimens becoming quite black. The cause of this change is not known.]

III. The Urine in Diseases of the Urinary Organs.

1. Renal Affections.

(a.) Acute Nephritis.—In this disease the urine is at first diminished in quantity—500–800 cc. or less being passed in twenty-four hours—of acid reaction, and high sp. gr. (1.015–1.025). The sp. gr. rarely attains to so great a height as in the urine of congestion. It ranges in colour from blood-red to that of a watery extract of meat, and the presence of blood-pigment in considerable quantity may be determined by *Heller's* test or by spectroscopic examination. In the latter case, if the urine be fresh, the characteristic bands of methæmoglobin may be visible.

Chemical analysis shows large quantities of albumin.

Microscopical investigation of the sediment affords the clue to the condition. It exhibits—

- (1.) Red blood-corpuscles in variable proportion. These are for the

most part altered, and present the appearance of washed-out discs (phantom corpuscles).

(2.) Some leucocytes. These are always less numerous than the phantom cells just mentioned.

(3.) Epithelium. Chiefly small polyhedral uninuclear cells from the urinary tubules, with a few others derived from the renal pelvis and bladder.

(4.) Casts. These are (a) formed of blood-corpuscles ; (b) formed of leucocytes ; (c) formed of renal epithelium ; (d) hyaline, more or less thickly beset with epithelial cells and red or white blood-corpuscles.

Such are the microscopical constituents of the sediment at the outset of an acute nephritis, as in the first and second days of the nephritis of scarlatina and erysipelas. They alter their character as the disease progresses, and after the lapse of a few days, and side by side with those described, appear the metamorphosed casts, granular and waxy, &c.

When in the course of chronic nephritis an acute exacerbation takes place, the urine possesses a similar character to that described above. But, as before, the description applies with full force only to the earlier period of the attack. If death from uræmia or œdema of the lungs does not ensue, the physical characters of the urine tend gradually to return to those of health. It is more abundant, the contained blood grows less, and then a light flesh-water tint alone declares the existence of acute nephritis. Albuminuria becomes less marked, and finally ceases with the approach of health. The other signs, recognisable only by means of the microscope, disappear with, or shortly after, the cessation of the albuminuria. It must be borne in mind that to justify the diagnosis of nephritis the formed material of the urine must be present in *considerable quantity*.

The occurrence of micro-organisms has been noticed in this chapter (see *Parasites*).

(b.) **Chronic Nephritis.**—The urine is normal in quantity, or somewhat lessened (1200–1500 cc. daily), acid, and of normal sp. gr. It usually contains a very considerable proportion of albumin. Microscopically the sediment is very variable in character, but renal epithelium is never absent, and the cells are often fatty. There are metamorphosed casts of different kinds, especially granular casts, and hyaline casts covered with blood-corpuscles or renal epithelium (p. 265). These are of special importance in diagnosis.

The occurrence of casts composed of fatty matter or overlaid with fat-crystals indicates advanced fatty degeneration of the renal tissue (p. 268).

It occasionally happens that, with all the symptoms of chronic nephritis present, no trace of casts or epithelium can be found in the urine. This occurs most often in long-standing and very chronic cases. *Schrwald*²¹ has called attention to the occasional absence of casts from the urine of nephritis, and believes that this is to be accounted for by their solution by pepsin (see p. 272) in acid urine. To guard against this, he suggests that the urine should be allowed to stand only for a short time and at a low temperature. To obtain the deposit by means of the sedimentator would undoubtedly add an element of certainty. *Glaser*²² has shown that, as the result of moderate indulgence in alcohol, the non-albuminous urine of healthy persons may contain abundance of leucocytes and casts of all kinds. These he ascribes to irritation of the kidney by alcohol. In conclusion, it may be stated that in rare cases of chronic nephritis the urine is altogether normal. *Stewart*²³ has repeatedly found the urine free from albumin in cases of chronic nephritis.

(c.) **Contracted Kidney.**—The quantity of urine is very greatly increased—4000 to 5000 cc. being passed in the twenty-four hours. Its reaction is acid and sp. gr. low (1.008–1.012 and less). In this respect, however, exceptions are not uncommon. The author has met with cases in which the quantity of the excretion was diminished and the sp. gr. proportionally increased. It is pale in colour, and contains but little albumin, sometimes only a mere trace. The sediment is usually scanty, and, when examined with the microscope, exhibits generally but a few hyaline and some granular casts.

It is to be observed that even those cases in which only a trace of albumin is to be found (small red kidney of *Ribbert*) often run a particularly unfavourable course.

(d.) **Amyloid Kidney.**—In this condition the urine presents characters which are for the most part similar to those of contracted kidney, that is to say, it is increased (though sometimes normal) in quantity, of acid reaction, and low sp. gr. As a rule, however, it contains more albumin.

Microscopical examination of the sediment shows hyaline casts in comparative abundance and some epithelium. The microscopical appearances, however, are subject to much variety, and the author has observed cases which were hardly to be distinguished from those of chronic nephritis. The amyloid reaction (iodo-potassic-iodide and sulphuric acid, &c.) with the casts cannot be depended upon, since, on the one hand, it is often obtained when the post-mortem appearances show no amyloid degeneration of the kidneys, and, on the other, it is absent in many cases where the symptoms (enlarged spleen and liver, &c.) point to this condition.

(e.) **Uræmia.**—The urine in cases where the symptoms of uræmia supervene is always rich in albumin, and microscopically resembles that of nephritis. It is scanty in a degree that sometimes amounts to anuria; and even when greatly reduced in quantity, we often find that

its sp. gr. is still below the healthy standard. Again, *the quantity excreted may remain unaltered, but the sp. gr. in such cases is always greatly reduced.*

When the urine is of this character, it would seem to hold poisonous bases in smaller quantity than ordinarily.⁶⁹⁴

What has been said here applies only to *typical forms* of renal disease. The appearances are modified according to the various anatomical perversions of the kidney structure upon which they depend.

Investigations which the author has made as to the character of the urine of children suffering from nephritis show that the important constituents of the excretion, urea, uric acid, sulphuric and phosphoric acid, are constantly diminished in quantity. In adults, according to *Munzer*, the urea is less, and the total elimination of nitrogen is always reduced. *v. Noorden* and *Ritter*⁶⁹⁵ observed a remarkable variation in this respect in renal disease. According to *Kornblum*,⁶⁹⁶ nitrogenous metabolism is much impeded in nephritis.

2. Pyelitis Calculosa.—The urine passed during the paroxysms of this affection is diminished in quantity, contains mucin in abundance, blood and pus in variable proportions, and concrete masses of uric acid or urates. After the paroxysms the urine is passed in great abundance. It is then pale in colour and of low sp. gr., and exhibits probably a flocculent precipitate of mucin. Persistent polyuria is common. When, as commonly happens, the condition is complicated with catarrh of the ureter and bladder, a more or less abundant purulent sediment forms independently of the attack.

According to *J. Fischl*,⁶⁹⁷ the urine at the commencement of this disease always contains casts, both hyaline and granular, and this authority regards their presence as of great importance in the differential diagnosis of pyelitis and cystitis. In addition, there occur plugs of cylindrical form composed of conglomerated white blood-corpuscles. These are probably derived from the renal pelvis, and indicate an extension of the processes to the kidney proper, or pyelonephritis. When the latter condition is established by the appearance of pyelitis, then the presence of nephritis, and granular casts, renal epithelium, &c., may be observed.

3. Ureteritis Membranacea.—In a case which was under the author's care, investigation of the urine disclosed a remarkable appearance. The patient was a woman with renal calculus causing pain. The urine held abundance of carbonate and sulphate of calcium and triple-phosphate, and in it were long spiral bodies of large size, which macroscopically, microscopically, and chemically exactly resembled *Curschmann's* spirals. There was no pus. The condition was probably an

affection of the ureter analogous to enteritis membranacea, and it might be appropriately called *ureteritis membranacea*.⁶⁹⁸ These membranous bodies continued to be discharged for some days.

4. Cystitis.—In uncomplicated cystitis the urine is generally pale, of normal sp. gr., and has an acid reaction, unless when alkaline fermentation takes place within the bladder. In the latter case it is turbid, and deposits on standing a more or less abundant deposit of fat-laden and swollen leucocytes and triple-phosphate crystals. Microscopically, moreover, pus cells and epithelium of various forms are to be seen. Amongst the latter, certain cells from the deeper layers of the mucous lining, provided with one or two finger-shaped processes, are especially noticeable. In connection with ichorous or hæmorrhagic cystitis, red blood-corpuscles and pigment masses appear in the urine. The complication of cystitis with an affection of the ureter cannot be determined by chemical and microscopical investigation, but its presence must be diagnosed from the existence of the other clinical symptoms.

*Schnitzler*⁶⁹⁹ has observed that the urine in cases of cystitis often contains a bacillus, pure cultivations of which, when transferred to the bladder of animals (rabbits), give rise to symptoms of the disease.⁷⁰⁰ *Escherich*⁷⁰¹ and others have shown that cystitis may be induced by the *Bacterium coli commune*.

Similar indications may be produced by a purulent urethritis, and confusion in this respect is to be guarded against.

The condition which has been designated as ammoniæmia, and which depends upon the absorption of ptomaines from the bladder, is often, but not always, accompanied with cystitis. In this connection the urine, when freshly voided, is undergoing alkaline fermentation (see p. 255).

5. Tuberculosis of the Urinary Organs.

(a.) **Tubercular Ulceration.**—Chemically and microscopically the urine in this condition resembles that of cystitis or pyelitis. It is pale, of normal sp. gr. and quantity, contains a variable proportion of albumin, and an abundant sediment, which consists principally of swollen and fat-laden pus-cells. The diagnosis must rest chiefly upon the recognition of tubercle-bacillus by the methods described in connection with the examination of the sputum (p. 126). In chronic tubercular affections the micro-organism is to be seen in great abundance in the urine, as in the specimen represented in fig. 114, and, as in this case, it tends to cohere in groups shaped like the letter S. The precise localisation of the affection must rest upon other grounds.

(b.) **Miliary Tuberculosis.**—In this affection the urine is often unchanged. It is apt, however, to contain blood at intervals, and may be distinguished from nephritis by the absence of casts and renal epi-

thelium. Tubercle bacilli are never to be found in considerable quantity in the sediment.⁷⁰²

6. Calculus and Tumours of the Bladder.—These are to be suspected when copious intermittent haemorrhages take place, and when the blood which is passed with the urine separates from it and is deposited in a thick layer at the bottom of the receiving vessel. The subjective symptoms, pain, &c., are usually sufficiently distinctive (see p. 290).

7. Catarrhal Urethritis.—The urine in catarrhal urethritis is altogether unchanged, except that the first flow contains pus. The affection is rarely met with. *Bockhart*⁷⁰³ ascribes it to infection with non-specific vaginal secretion.

8. Gonorrhœal Urethritis.—The appearances are the same as in simple urethritis, but pus is usually very copious. It would appear that in all cases, while the infection is recent, specific micro-organisms, gonococci, are to be found. These were discovered by *Neisser*. They are diminutive roll-shaped cocci, aggregated in large groups, which often closely pack the exfoliated epithelium cells of the urethra and cover their surface (*Neisser, Bumm, Bockhart*).⁷⁰⁴ More recent experience tended at first to lessen the clinical significance of these forms (*v. Zeissl, Hartdegen, Wendt*),⁷⁰⁵ since bodies altogether resembling gonococci have been found to be present in the genital tract under the most dissimilar conditions. The researches of *Wertheim*⁷⁰⁶ in Schauta's clinic, however, have quite lately established their specific character beyond any doubt. According to *Roux*,⁷⁰⁷ the supposed gonorrhœal microbe may be distinguished from other forms by the fact that it does not stain with *Gram's* method. *C. Schutz's*⁷⁰⁸ process is as follows:—The prepared cover-glasses are placed in a semi-saturated solution of methylene-blue holding 5 per cent. carbolic acid, and left in it for 5–10 minutes. They are then removed and washed in distilled water, to which acetic acid has been added in the proportion of five drops of the dilute acid to 20 cc. of water. After this they are given the contrast-stain in a very dilute solution of safranin. This method furnishes good specimens, but there is no reason to prefer it to staining with carbol-fuchsin, by which the specimens figured here were obtained. Fig. 142 represents gonorrhœal pus from a case of old infection; fig. 143 shows gonococci from a preparation of *Dr. Kolisko*, made with the pus two days after the infecting coitus.⁷⁰⁹

{ *McCann*⁷¹⁰ recommends as a cultivation medium for the gonococcus, the fluid taken with aseptic precautions from an ovarian cyst } *Wertheim*⁷¹¹ employs peptone agar bouillon seasoned with human blood, and in *Sternschneider's*⁷¹² experience the addition to the nutrient substance of sterilised urine promotes exuberant development. The gonococci produce on such media transparent dewdrop-like cultures. They fail to propagate on agar, and this fact is available for diagnosis.

The occurrence of gonococci threads and hyaline epithelium occurring in the urinary sediment of this condition is worthy of notice.⁷¹³

IV. The Urine in Diseases of the Alimentary Canal.—Diseases of the alimentary canal, for the most part, do not specially affect the urine pathologically; but in general it may be stated, that where they are attended with increased albuminous decomposition within the intestine, that fluid is apt to contain a large amount of indican. Carcinoma of the stomach with ulceration is occasionally attested by the appearance of peptone in considerable quantities (*Maienner*). In chronic catarrh of the stomach, and especially in dyspepsia, the acidity of the urine is apt to be much lessened.

V. The Urine in Hepatic Affections.—It may be stated in general that in all diseases which seriously involve the proper struc-

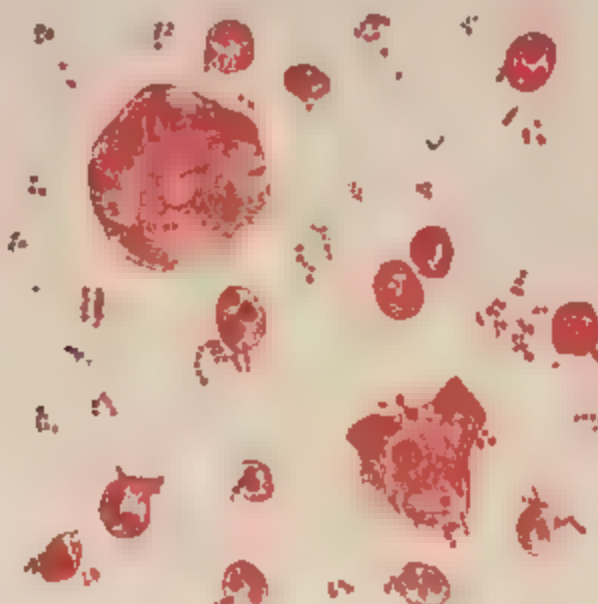


FIG. 142.—Gonococci of Urethritis (eye-piece III., objective, oil-immersion $\frac{1}{3}$, *Zeiss*).

ture of the liver, the excretion of urea is diminished, and even in certain severe forms (acute yellow atrophy) entirely suppressed (*Schultzen* and *Rieß*).⁷¹⁴ It is then to some extent represented by other nitrogenous metabolites which appear in the urine as leucin and tyrosin⁷¹⁵ (comp. p. 285), and together with these certain non-nitrogenous substances. Oxyamygdalic acid (*Schultzen* and *Rieß*, *Röhmann*⁷¹⁶), lactic acid, and the volatile fatty acids are often eliminated, as in cancer and syphilis affecting the liver (*r. Jakob*).⁷¹⁷

Diseases which cause an obstruction to the flow of bile are attended with the presence of bile pigments in the urine (see p. 336).

In atrophic cirrhosis the latter is scanty, rich in urates, and almost or entirely devoid of biliary colouring-matters, but often contains a high percentage of urobilin. In hypertrophic cirrhosis it is often quite

normal in quantity, sometimes increased, and exhibits abundance of bile-pigment.

The appearance of sugar and albumin in connection with disease of the liver is too uncertain to be of service in diagnosis. According to *Kraus* and *Ludwig*,⁷¹⁸ the ingestion of carbohydrates (grape-sugar) when the liver is diseased is attended by glycosuria. This seems to imply a profound change in the structure of the hepatic cells, and alimentary glycosuria is to be met with in atrophic affections of the liver of whatever kind. In cirrhosis the urine often holds grape-sugar in small quantity.⁷¹⁹ The chemical constitution of the urine in hepatic disease generally exhibits much variety.⁷²⁰ *Luigi Bellati*⁷²¹ has observed that its toxic properties are increased.

VI. The Urine in Diabetes Mellitus.—In this disease the urine

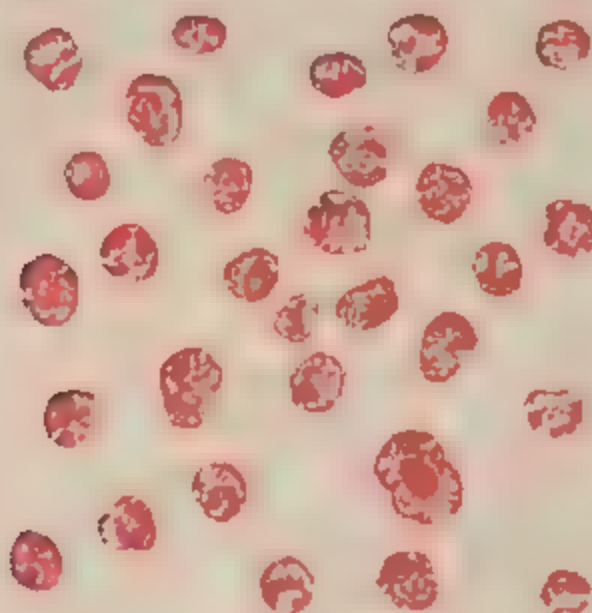


FIG. 143. Gonococci (two days after infection).

is pale and clear, and in colour often inclines to green. Its quantity is enormously increased, as much as 12-15 litres being passed in twenty-four hours, sp. gr. high, ranging from 1.030-1.050. It is usually rich in the indigo-forming substance, and invariably contains a greater or less proportion of grape-sugar (p. 317). Towards the close of the disease albumin in considerable quantity is apt to make its appearance (*Stohris*).⁷²² Exceptional cases of diabetes are met with in which the quantity of urine is not increased, and its sp. gr. is even lowered. When diabetes is complicated by acute disease, the excretion of sugar may entirely cease, as has been pointed out by *R. v. Engel*⁷²³ in a contribution from the author's clinic. In certain mild cases of diabetes, glycosuria occurs only after the ingestion of carbohydrates in large quantities.

The author has met with one case of diabetes mellitus, in the practice of Professor Nothnagel, in which the urine contained much acetone, and showed a sp. gr. of 1.003. It was also found to hold more than 0.3 per cent. sugar in solution.

Other substances that are occasionally to be found in the urine of diabetes are acetone in large quantity, diacetic acid, and a number of other organic acids, such as β -oxy-butyric acid,⁷²⁴ fatty acids,⁷²⁵ &c. [The urea of diabetic urine is in excess of the normal—uric acid is unaffected or diminished (*Taylor*). Ammonia, according to *Stadelmann*, is present in large quantity, but it is neutralised and the urine remains acid in consequence of the β -oxy-butyric acid which it contains.]

In view of the fact that β -oxy-butyric acid occurs in the urine of diabetes, and of other conditions, as febrile states, some notice of the method of detecting that body is called for. The following method by *Külv* is the best.⁷²⁶ The grape-sugar in the urine is fermented with yeast, the fluid filtered, and the filtrate concentrated to a syrupy consistence. The latter is mixed with its own bulk of concentrated sulphuric acid, the mixture distilled, and the distillate collected in a test-glass. If β -oxy-butyric acid be present, crystals of α -crotonic acid separate on cooling from the contents of the test-glass, and may be recognised by determination of their melting-point (72° C.). Should no such crystals form in the process, the distillate is shaken up with æther, and the crystals which deposit on evaporation of the æther are tested to ascertain their melting-point.

[The researches of *Stadelmann* and others point to β -oxy-butyric acid as the toxic agent in diabetes. It yields diacetic acid on oxidation, and this breaks up into CO₂ and acetone.]

Diabetic urine contains other carbohydrates, fruit-sugar,⁷²⁷ dextrin, &c. Albuminuria is apt to arise, and in a series of thirty-two cases examined, albumin was found thirteen times (*r. Jaksch*). The occurrence of coma is almost always attended by the presence of albumin, and the quantity of sugar at the same time diminishes. According to *Külv*,⁷²⁸ casts occur at the commencement of diabetic coma. In the author's experience the casts are surprisingly numerous in this condition. They are insoluble in æther and chloroform, colour mahogany brown with iodo-potassic-iodide solution, and acquire a light-blue tint on the application of sulphuric acid.

VII. The Urine in Diabetes Insipidus.—Marked polyuria is the characteristic symptom. Sixteen to twenty litres may be passed in the day. The urine is clear and almost colourless, of greatly reduced sp. gr. (1.0001-1.004). It contains neither albumin nor sugar, although it sometimes yields a small proportion of indican and inosite.

In a typical case of diabetes insipidus the nitrogen constituent of the fæces was remarkably small.

VIII. The Urine in Anæmia.—The fluid is pale, of low sp. gr., and neutral or alkaline in reaction. [Though pale when passed, it sometimes turns a deep red colour on the addition of nitric acid (*Taylor*). This depends on the presence of a chromogen (see above).] In the later stages of a severe anæmia albumin may be found, whilst at the same time, except for a few hyaline casts, tissue elements are entirely absent from the sediment (*Bamberger's hæmatogenic albuminuria*).

[The urine of pernicious anæmia is of low sp. gr. and very highly coloured. It contains (1) pathological urobilin; (2) granules of blood pigment, microscopically; (3) excess of iron. These characters are of much importance in diagnosis (*Hunter*).⁷²⁹

According to *Hunter*, the elimination of iron by the urine, while diminished in chlorosis, is greatly increased in pernicious anæmia. He estimates the daily output as follows: In health 5.6 mgrms.; in chlorosis, 1.7 mgrm.; in pernicious anæmia, 32.2 mgrms. The same observer has found diamines in the urine of this disease.⁷³⁰]

In leukæmia the elimination of uric acid is increased.⁷³¹ *Prus*⁷³² has observed leucin, and lactic acid is also found in the urine.⁷³³ The urine is rich in nucleo-albumin,⁷³⁴ but seldom contains peptone.⁷³⁵ Histone has been found by *Kolisch* and *Burian*.⁷³⁶

It will be convenient here to describe the method by which the lactic acid may be detected.

Detection of Lactic Acid.—*Schütz's* process:⁷³⁷—The urine passed in twenty-four hours is collected, and a precipitate thrown down with neutral acetate of lead. To the filtrate sulphuretted hydrogen is added, and it is then evaporated to a syrupy consistence, boiled with alcohol, filtered, and the alcohol distilled or driven off on the water-bath. The residue is freely treated with phosphoric acid and extracted for twenty-four hours in Schwartz's apparatus (see p. 85). The dark-brown oily fluid in the receiver is freed from æther by evaporation. The residue is dissolved in water, boiled with excess of zinc carbonate, and filtered. The remaining zinc carbonate is repeatedly boiled with water, the filtrates collected, concentrated on the water-bath, treated with 69 per cent. alcohol, filtered, and to the filtrate æther is added till a precipitate ceases to form. The mixture is allowed to stand, and the resulting crystals inspected. Their appearance—small prisms—and content, as shown by analysis of water and of zinc, will serve to identify the zinc salt of lactic acid.

Colasanti and *Moscatelli*⁷³⁸ assert that sarcolactic acid appears in the urine normally after great bodily exertion. *Heuss*⁷³⁹ failed to detect lactic acid in the urine either of health or in osteomalacia. *Schütz*⁷⁴⁰ failed to find lactic acid in hepatic disease, pernicious anæmia, and leukæmia. *Munzer* and *Palma*⁷⁴¹ determined the presence of the acid in carbonic oxide poisoning.

IX. The Urine of Toxic States.

1. Poisoning with Acids.—Poisoning with strong mineral acids⁷⁴²—nitric, sulphuric, and hydrochloric—is generally followed by the appearance of blood and albumin in the urine. The symptoms, however, are often very transient. Toxic nephritis may ensue, and this especially when sulphuric acid was the poison. The urine is then scanty, of high sp. gr. and acid reaction. Chemically and microscopically, it presents the characters of that of acute nephritis (see above). In every case of poisoning with acids which the author has had an opportunity of observing, the urine possessed the property of dissolving cupric sulphate in alkaline solution, and reducing it when subsequently boiled, whilst at the same time the most sensitive tests besides gave no indication of sugar.

2. Poisoning with Alkalies.—After poisoning with caustic potash, which may be taken as the type of this condition, the urine passed during the first few hours contains albumin, invariably in severe cases, but very often also when the symptoms are slight. Its chemical and microscopical characters at the same time are not otherwise those of nephritis. Its reaction is feebly acid, seldom neutral, very rarely alkaline. It possesses the reducing property in a marked degree, while no evidence of sugar can be obtained with the phenylhydrazin reaction and other tests. Where the toxic agent has been chlorate of potash, acute nephritis is apt to follow. The detection of this salt in the urine may be effected by the method described at p. 181.

3. Poisoning with Metals and Metalloids.

(a.) **Lead Salts.**—In acute lead-poisoning, especially when attended with colic, large quantities of albumin are often transitorily present in the urine; more frequently still a true renal albuminuria, due to secondary nephritis, occurs. The presence of lead may be determined directly by the method described in connection with the vomit (see p. 181).

(b.) **Salts of Mercury.**—The urine in mercurial poisoning, after the lapse of a few hours, contains a large proportion of albumin, and very often blood. The symptoms of nephritis are usually sooner or later developed. This is especially true of poisoning with corrosive sublimate.⁷⁴³

Mercury may be detected in the same manner as in the vomit (p. 182), or in the urine *Fürbringer's* process may be employed,⁷⁴⁴ or, better still, *Ludwig's* method.⁷⁴⁵

Five hundred cc. of urine are acidified with hydrochloric acid (1-2 cc.) in a beaker and heated to 50-60° C. Three grms. of granular zinc or finely divided copper are then placed in the fluid, which is shaken up for half a minute. The metal is then allowed to settle, the supernatant fluid poured off, and the sedi-

ment obtained upon a filter, where it is well washed with boiling water, and dried at 60° C. The powdered metal is then placed in a tube of hard glass, of 8 to mm diameter, closed at one end, covered with a plug of asbestos, upon which again is put a layer some 5-6 cc deep of granular oxide of copper; then another asbestos plug, and, finally, another layer of granular zinc, which has been previously dried and well heated. When the tube is filled in this way, it is drawn out to capillary calibre at a point some millimetres behind the last asbestos plug, and a bulbous expansion is made at the end. The cupric oxide is first heated to a dull red, the layer of zinc to a lesser degree, and, finally, the powdered metal containing mercury.

The mercury is deposited as a metallic powder in the capillary tube. The latter is then broken off above the last of the asbestos rings by letting water drop on it; a few particles of metallic iodine are placed in the first part of the tube while still hot, and the other expanded end of the capillary is connected with an aspirator (Bohm's air-pump serves best). The iodine vapour impinges on the mercury, and mercuric iodide forms, and is easily distinguished by its colour.⁷⁴⁶

Wolf and *Nega*⁷⁴⁷ have employed a modification of this process. Organic matter is first removed by hydrochloric acid and chlorate of potash, and copper as a thin foil is introduced to receive the metal (mercury) instead of granular zinc or powdered copper. The method is said to be very satisfactory. That of *Alt*⁷⁴⁸ is still more simple. *Winternitz*⁷⁴⁹ has devised a method for the quantitative estimation of mercury in the urine.

*Almen's*⁷⁵⁰ method is the following.—About 300 cc. of the urine to be tested are taken, a little caustic soda and some sugar added, and the mixture boiled. The phosphatic sediment which falls carries the mercury with it. When it has entirely settled, the fluid is decanted off, the sediment dissolved in hydrochloric acid, and a piece of fine copper or brass wire placed in the fluid, which is then maintained at a moderate heat for an hour and a half. After this the wire is removed, boiled in alkaline water, and dried with blotting-paper. It is then placed in a glass tube of small calibre, which is broken a few millimetres in front of the wire, fused at the end, and heated over a small flame. The mercury sublimes, and is deposited in small globules, which can be readily recognised with the microscope. The reagents employed in the process should be previously tested for such mercury as they may themselves contain (see p. 182).

(c.) **Salts of Copper.**—In poisoning with copper salts, the urine is always reduced in quantity. It usually contains albumin, and very often also blood. As to whether acute nephritis can originate in this way, some doubt exists. Experiments on animals would seem to favour the assumption. To detect the poison in the urine, the method described at p. 183 will serve.

(d.) **Arsenic.**—In acute arsenical poisoning albumin and sometimes blood in considerable quantities appear in the urine. In one case the author has observed all the signs of acute nephritis. The urine possesses the reducing property, but sugar cannot otherwise be made evident. The effect on the urine of chronic arsenic-poisoning is but little understood. Albuminuria would seem occasionally to occur in this connection.

For the detection of arsenic, see chapter on the vomit (p. 183).

(e.) **Phosphorus.**—At the outset of a case of phosphorus-poisoning the urine is not notably changed as to quantity or sp. gr. Later it contains albumin, but not usually in considerable quantity, occasionally blood, and very commonly casts of different kinds. Hyaline casts, with bile-stained lining cells of the renal tubules adhering to them, are seen, and also casts formed of renal cells that have undergone fatty degeneration, but only at a late period of the disease. Leucin and tyrosin are seldom present, and in some instances where they have been described it is probable that the substances observed were really composed of the lime and magnesia salts of the higher fatty acids, the formation of which in various morbid states was pointed out by the author some years ago.⁷⁵¹ Hæmatoidin crystals are met with. *E. Schütz*⁷⁵² has recorded a case in which large quantities of fat were voided. Sugar and carbohydrates are not as a rule eliminated in any large amount, but the author has shown that the typical phosphorus-liver leads to alimentary glycosuria. *Maixner* and the author⁷⁵³ have found peptone in a few cases, and according to *W. Robitschek*⁷⁵⁴ a transitory peptonuria is a common incident in phosphorus-poisoning. *Münzer*,⁷⁵⁵ working in the author's clinic, has ascertained that in the earliest stage of this condition the excretion of nitrogen is lessened and quickly falls to a very low figure. *O. Storch*⁷⁵⁶ obtained similar results from experiments on animals so long ago as the year 1865. Uric acid is but little diminished, and consequently the change is chiefly in respect of urea. There is a remarkable formation of ammonia, which serves to neutralise the great excess of acidity due to acid products of the decomposition of proteids. Chlorides are scanty. Phosphorus is as early as the second day increased in proportion to the nitrogenous increase, and to the extent of 50 or even 90 per cent., then diminishes until death, or in favourable cases the amount rises again as convalescence approaches. This is true also of sulphuric acid. Fatty (*v. Jaksch*) and lactic (*Schultzen-Riess* and *Münzer*) acids are also excreted.⁷⁵⁷

4. Poisoning with Alkaloids.

(a.) **Morphia.**—In acute poisoning with morphia the urine is commonly found to contain sugar. In chronic morphinism, again, it has a powerful reducing action, and sugar may sometimes be found with other tests (see p. 317). [The reducing property depends upon the presence of glycuronic acid (*Halliburton*).] For the detection of morphia in the urine the method described at p. 185 may be adopted. It must be borne in mind, however, that morphia does not appear in the urine in all cases of morphia poisoning or morphinism, and its absence does not prove that the alkaloid has not been taken. The researches of *Donath*⁷⁵⁸ have shown that it may disappear entirely within the system.

(b.) **Nicotin.**—The toxic effects of nicotin are not associated with any special changes in the character of the urine. For the detection of the alkaloid see p. 186.

(c.) **Atropin.**—Little that is definite is known of this condition.

Atropin may be isolated from the urine in the same manner as that described in connection with the vomit (comp. Chap. IV.). As a test for its presence a little of the urine may be placed in an animal's eye, and the mydriatic effect watched for. This will be obtained, according to *de Ruitter and Donders*,⁷⁵⁹ when the urine contains the alkaloid in the proportion of one part to 130,000 of water. In cases of poisoning with deadly nightshade berries (*Atropa belladonna*), the urine has a peculiar fluorescence (*A. Paltans*,⁷⁶⁰) due to the presence of scopolitin. This does not occur in poisoning with the pure alkaloid, and in certain circumstances it affords an indication as to the source of the poison.

(d.) **Ptomaines (Exogenic Toxicosis).** The phenomena of poisoning with ptomaines need further investigation. In a case (of sausage-poisoning) which was recently under the care of Professor Nethnagel, albuminuria and the signs of nephritis supervened. Subsequent experience has convinced the author that kidney trouble regularly occurs in the later stages of ptomaine-poisoning.

5. Poisoning with Ethylic Alcohol.—Chronic alcohol-poisoning appears to produce nephritis and arterio-sclerosis.⁷⁶¹ In the acute condition the poison can be found only as the merest trace in the urine.⁷⁶² To recognise its presence the urine must be distilled by means of a steam bath, and the distillate treated in the manner described at p. 189.

6. Chloroform Poisoning. In this condition the urine is generally of high sp. gr. It often contains a trace of albumin and some sugar [or glycuronic acid (*Halliburton*)]⁷⁶³ According to *Kast and Meister*,⁷⁶⁴ the urine after the prolonged administration of chloroform contains an organic sulphur compound and urobilin, and is highly toxic.⁷⁶⁵ For the detection of chloroform the urine is distilled by means of a steam-bath to prevent frothing, and the first drops of the distillate are submitted to Hoffman's or Vitali's test. The results obtained in this way are equally useful with those of *Marichal's* method.⁷⁶⁶

7. Carbolic Acid Poisoning. When large quantities of carbolic acid have been administered through the mouth or absorbed from a wound, the voided urine assumes a dark green colour, which changes to black on standing. This colour is due to the presence of hydrochinon, a derivative of phenol, which, while still within the system, is in part transformed into coloured oxidation products (*Baumann and Preuse*).⁷⁶⁷ Pure phenol is never present, but even in the severest cases is found only in combination with sulphuric acid (p. 346). Hence the characteristic reaction of carbolic acid—a violet coloration with

solution of perchloride of iron—cannot be obtained with the urine. The latter contains a small quantity of albumin, and commonly also hæmoglobin. A further evidence of the condition depends upon the diminished proportion of simple sulphuric acid which it contains. When chloride of barium is added to healthy urine acidulated with acetic acid, an abundant precipitate of barium sulphate falls ; but when from any cause the quantity of sulphuric acid in the uncombined form is lessened, this precipitate fails, or is represented by a mere turbidity. If now in this case the urine be filtered and boiled with hydrochloric acid, so as to decompose the phenol-sulphuric acid, with the reproduction of simple sulphuric acid a copious precipitate of barium sulphate forms.

Inasmuch as phenol-sulphuric acid is normally a constituent of the urine, it serves but little purpose in cases of carbolic acid poisoning to attempt an estimate of the phenol which passes over as tribromo-phenol in the process of distillation. A better plan is to determine the relative proportion of simple and compound sulphuric acid present. If it be found that the latter is increased while the first is diminished, and if at the same time such affections as promote the elimination of æther-sulphuric acids (active albuminous decomposition) can be excluded, this is strong presumptive evidence of carbolic acid poisoning.

The same method will serve for the detection of all the aromatic substances which appear in the urine as æther-sulphuric acids in cases of poisoning or after the administration of drugs.

8. Poisoning with Nitro-Benzol and Aniline.

(a.) **Nitro-Benzol**.—In cases of poisoning with this substance, the urine has the odour of nitro-benzol, and generally contains a substance which has the property of rotating polarised light to the left, and of reducing cupric sulphate in alkaline solution.⁷⁰⁸ There is a slight increase in the amount of ammonia and of acetone excreted (*Münzer* and *Palma*,⁷⁰⁹ *Bonde*⁷⁷⁰). Traces of sugar are found, and *Strasser* has obtained alimentary glycosuria.

(b.) **Aniline**.—The recorded observations of cases of poisoning with aniline show that the character of the urine varies considerably. It is usually dark in colour and highly concentrated (*Grandhomme*).⁷⁷¹ In one instance (*Fr. Müller*⁷⁷²) it was free from sugar, albumin, and blood, and exhibited the reduction phenomenon in a marked degree. Æther-sulphuric acids were notably increased. An æthereal extract was found to contain aniline by its yielding a violet colour in presence of solution of chloride of lime. *Müller* is of opinion that this substance is in part eliminated as para-amido-phenol-sulphuric acid.

9. **Poisoning with Carbonic Oxide Gas.**—The urine passed in this condition invariably contains grape-sugar⁷⁷³ (see p. 316) and an

uncertain proportion of albumin. The quantity of sugar varies directly with the intensity of the poisoning, and is increased by administration of carbohydrates. According to *Munzer* and *Palma*,⁷⁴ the output of uric acid is much greater than normal; that of ammonia and acetone but little increased. Lactic acid is present.

V. THE DETECTION OF CERTAIN DRUGS IN THE URINE.

1. Iodoform and Salts of Iodine and Bromine.—After the exhibition of iodoform, whether internally or by outward application, iodides and iodates may be detected in the urine. So too with iodine when applied externally as the tincture, or taken by the mouth as iodide of potassium, it may be readily recognised in that fluid, and the administration of thyroid extract has the same effect.⁷⁵

To test for iodine : (a.) the urine is treated with a little fuming nitric acid or chlorine water, and shaken up with chloroform. If a salt of iodine be present, the metal is set free, and dissolves in the chloroform, with the formation of a red colour. (b.) *Sandland*⁷⁶ treats the urine with dilute sulphuric acid and sodium nitrite, and separates with bisulphide of carbon. (c.) *Jolles'* test :⁷⁷—Concentrated hydrochloric acid is added in equal quantity to the urine, and on the surface of the fluid a little chloride of lime solution is poured. At the line of junction there forms a brown ring which turns blue on the addition of starch solution.

The quantitative estimation may be effected best by *Harnack's*⁷⁸ method, in which all the iodine is converted into palladium iodide.⁷⁹ Iodine appears in the urine very shortly after it has been taken into the system. A quarter of an hour will suffice for its manifestation there.

Bromine Salts, when very abundant, may be detected thus:—The urine is treated with chlorine water, and shaken up with chloroform, when bromine dissolves in the latter, with the production of a yellow colour. It will usually be found necessary, however, to evaporate the urine previously, then to incinerate it carefully, and to test the colourless watery extract from the ash in the manner described above.

2 Salicylates - Salol and Betol.—Salicylates also very quickly pass into the urine. The latter then exhibits a remarkable reducing power, and yields a violet colour in presence of solution of perchloride of iron. This reaction depends partly upon the presence of salicylic acid and partly upon that of salicyluric acid, into which the latter is changed within the system. This reaction is not readily prevented by boiling. The body on which it depends is taken up by æther from acidified urine, and can be detected in the æthereal extract by means of solution of perchloride of iron. The reaction, unlike that of diacetic

acid (see *Diaceturia*), does not disappear on standing. It is well in testing for it to precipitate the phosphates with solution of perchloride of iron, and filter, and then test the filtrate with more of the reagent.

The exhibition of salol (phenyl-æther of salicylic acid) imparts a similar character to the urine, which also on standing acquires a tint varying from dark green to black, like that following the use of carbolic acid.

. The use of betol (naphthalol, salicylate of β -naphthol-æther) does not impart any special colour to the urine, which, however, yields the perchloride of iron reaction. Both salol and betol appear in the urine as sulphuric acid combinations (*v. Jaksch*).⁷⁸⁰

3. Quinine, Kairin, Antipyrin, Thallin, Antifebrin, Phenacetin, and Lactophenin.

(a.) **Quinine.**—This alkaloid is said by *Kerner*⁷⁸¹ to be eliminated as dioxyquinin. It renders the urine dark. For its detection a large quantity of the fluid is taken, treated with ammonia, and shaken up with æther. The latter is then distilled off or evaporated, and quinine remains in the residue. It is dissolved in acidulated water, and the addition of chlorine water and ammonia causes an emerald-green tint to develop.

(b.) **Kairin.**—Under the influence of this drug the urine acquires a brown colour, which becomes brownish red with solution of perchloride of iron. The substance upon which this reaction depends may be extracted with æther from acidified urine. The reaction in the æthereal extract remains even after the lapse of weeks. The addition of strong acids to the urine destroys the reaction, and it is, moreover, impaired by prolonged boiling. According to *v. Mering*,⁷⁸² kairin is eliminated as kairin-potassium sulphate.

(c.) **Antipyrin** imparts a darker tint to the urine, which gradually becomes a purple-red when treated with perchloride of iron. If the urine be acidulated and extracted with æther, a substance is obtained which colours brown with perchloride of iron. This reaction is gradually lost after the lapse of a few days. It is impaired, but not destroyed, by boiling the urine. It is destroyed by the addition of acids. Estimation of the quantities of simple and of æther-sulphuric acids in appropriate instances has shown that antipyrin is eliminated as a sulphuric acid combination (*v. Jaksch*).

(d.) **Thallin.**—The urine after the administration of thallin is usually a brownish green when viewed in bulk, greenish in a thin layer. Treated with perchloride of iron, it presently exhibits a purple-red tint, which, after the lapse of four or five hours, if undisturbed, passes into a brownish red. If a little mineral acid be added and the fluid be shaken up with æther, the latter takes up a substance which colours

brown-red with perchloride of iron. The coloration does not vanish when the specimen is allowed to stand, but rather becomes more marked. When the pure thallin urine is shaken up with æther, the æthereal extract contains a body which colours green with perchloride of iron (thallin).⁷⁸³ The tint in this case disappears on prolonged standing. The red colour obtained with perchloride is lost by boiling for a few seconds. It is likewise destroyed by mineral acids. Thallin is partly eliminated in the form of chinanisol.

(e.) **Antifebrin.**—The physical character of the urine is unaffected even by large doses of antifebrin. Chemically it has been ascertained by *F. Müller* that it contains more than the normal proportion of æther-sulphuric acids; and this is accounted for by the supposition that antifebrin is changed within the system to paraamidophenol-sulphuric acid. Where the drug has been exhibited in large quantities, it may be recognised in the urine (*F. Müller*⁷⁸⁴) thus:—The fluid is boiled with one-fourth its bulk of strong hydrochloric acid, and when it has cooled, a few cc. of a 3 per cent. solution of carbolic acid is added, and then a few drops of a solution of chromic acid. If paraamidophenol be present, the specimen develops a red colour, which gives place to blue on the addition of ammonia.

For the detection of antifebrin *Yron*⁷⁸⁵ recommends the following method:—The urine is shaken up with chloroform, and the residue from the extract is heated with a little mercurous nitrate. If antifebrin be present an intense green coloration takes place. When antifebrin has been separated from the urine, as by shaking it up with æther and acid, it can also be recognised by the addition of chloroform and caustic potash. Antifebrin is excreted for the most part in combination with sulphuric acid (*Fr. Müller, Mörner, v. Jaksch*). According to *Mörner*,⁷⁸⁶ this body is in part oxidised to acetylparaamidophenol within the system, and eliminated as æther-sulphuric acid.

(f.) **Phenacetin (acetphenetidin).**—The colour of the urine is unaffected by this body, even in large doses. It rotates polarised light to the left (glycuronic acid combination, *Fr. Müller*⁷⁸⁷), and exhibits the paraamidophenol reaction described above. It contains no free phenacetin, but the presence of phenetidin may be shown by changing it into its diazo compound, which then yields a purple coloration with naphthol or yellow with phenol (*Müller*). The process is as follows:—To a specimen of the urine two drops each of hydrochloric acid and of sodium nitrite solution (1 per cent.) are added. The further addition of an alkaline watery solution of *a*-naphthol and a little caustic soda causes a beautiful red colour to develop, and this passes into violet if hydrochloric acid be supplied. Under the same conditions phenol gives a citron-yellow in alkaline and a rose colour in acid solution. When

large doses of the drug have been taken, the urine gradually takes a brown-red tint in presence of perchloride of iron solutions and oxidising substances, changing slowly to black on standing for a long time. According to *Ubaldi*,⁷⁸⁸ the ingestion of phenacetin is attended by an increase in the amount of combined sulphuric acid in the urine.

(g.) **Lactophenin.**—This substance turns the urine dark, and the colour deepens on standing. It gives the paraamidophenol reaction. No change occurs on the addition of ferric chloride.⁷⁸⁹

4. Chrysophanic Acid.—The administration of infusion of senna or of preparations of rhubarb imparts a reddish-brown colour to the urine, which may be present in the freshly voided fluid or develop on standing. This gives place to red on the addition of alkalies at ordinary temperatures. When boiled with alkalies the resulting phosphatic sediment is not red, but yellow; and if dissolved in acetic acid the solution also colours yellow, changing to violet on exposure to the air; in this respect differing from the precipitate of blood pigments, which is also soluble in acetic acid, but gradually bleaches when exposed.

5. Santonin.—Santonin taken internally colours the urine yellow, which again gives place to red on the addition of alkalies. Its presence may be discriminated from that of rhubarb (*Munk*)⁷⁹⁰ from the fact that the red colour developed by the latter in presence of alkalies, though permanent, is rapidly destroyed by reducing agents (granular zinc, sodium amalgam), whilst that due to santonin persists under like circumstances. Chrysophanic acid is precipitated with baryta water. The precipitate is red, and leaves a colourless filtrate. The latter is yellow if santonin be present. The addition of alkaline carbonates turns the urine red, rapidly if it contain rhubarb, very slowly and gradually in the case of santonin.

G. Hoppe-Seyler⁷⁹¹ has suggested the following method for the discrimination of chrysophanic acid and santonin:—The urine is treated with caustic soda and extracted with amylic alcohol. If the colouring matter of santonin be present, it passes over with the alcohol, and the specimen is decolorised. The colouring matter of chrysophanic acid derived from the exhibition of rhubarb or of senna is little or not at all taken up by amylic alcohol in presence of the alkali.

6. Tannin.—When tannin has been taken medicinally in large doses, the urine turns dark green with solution of perchloride of iron.

7. Naphthalin.—Naphthalin in large doses causes the urine to assume, especially on standing for a long time, a dark tint, like that due to carbolic acid.

According to *Penzoldt*,⁷⁹² a dark-green colour develops rapidly in presence of concentrated sulphuric acid.

8. Copaiba Balsam.—Copaiba balsam in the urine yields a red

colour with hydrochloric acid, changing to violet when heated. If ammonia or caustic soda be added to the urine, a light-brown colour with a blue fluorescence develops (*Edlefsen*).⁷⁹³

When boiled with an acid the urine gives a precipitate which is soluble in alcohol.

It may be mentioned here that after the use of oil of turpentine the urine sometimes gives a precipitate with acids. It has also a characteristic odour of violets.

Myrtol imparts a similar odour, and when the urine is superimposed upon nitric acid a red ring gradually develops.

CHAPTER VIII

INVESTIGATION OF EXUDATIONS, TRANSUDATIONS, AND CYSTIC FLUIDS

FLUID may be effused into any of the cavities of the body as a consequence of inflammation or of disturbances in the circulatory system.

Under appropriate circumstances a portion of such fluid may be drawn off by puncture or in some other way, or a spontaneous opening may occur, and in either case the fluids so obtained may be submitted to examination. In this way much useful information may be secured for the purposes of diagnosis. A question arises at the outset as to whether the fluid is an inflammatory product (exudation), or the result of impediment to the circulation, or derived from the degeneration of certain organs (transudation).

A.—EXUDATIONS.

An exudation may be purulent, sero-purulent, putrid, hæmorrhagic, or serous. All such fluids, with the exception of the last two, imply, of course, an inflammatory origin. Upon its other characters, and especially upon the nature of the tissue elements which it contains, more precise inferences may be based in the case of any one of them.

i. PURULENT EXUDATIONS.

I. NAKED-EYE APPEARANCES.—Pus (*bonum et laudabile*) is a turbid fluid of varying consistence and high sp. gr., with an alkaline reaction, and ranging in colour from grey to a greenish yellow. It may accumulate in natural cavities (exudations), or be effused amongst the tissues (phlegmon), or, finally, it may be secreted from the surface of a wound. On standing in a cool place it separates into two layers, the upper of which is of a light-yellow colour and tolerably transparent, and the lower opaque from the deposit of pus-cells. It is often brown or brownish red from admixture with blood. Putrid pus can always be discriminated by its naked-eye properties. It is thin, green, or brownish red, and emits an extremely penetrating odour of indol and skatol.

It is apt, especially when derived from the intestinal tract, to contain sulphuretted hydrogen, which may be recognised by the sense of smell. The gas may find its way from the alimentary canal without any obvious channel of communication, or may be generated independently through the agency of micro-organisms. The chemical determination of this gas has been dealt with elsewhere.

II. MICROSCOPICAL CHARACTERS.

1. White and Red Blood-Corpuscles and Epithelium.—When pus is submitted to microscopical examination, it discloses a multitude of cells which entirely resemble white blood-corpuscles. When derived from a perfectly fresh specimen, these cells exhibit the contractile property, and the mahogany coloration obtained with solution of iodine and iodide of potassium or ammonium attests the presence of glycogen. This reaction is obtained best with fresh pus from the surface of a wound. The dead cells are shrunken and very granular, or may appear as decomposed or decomposing particles of protoplasm.

In addition to this, giant pus-corpuscles and fat-laden cells are occasionally observed. No special significance attaches to their presence. They have been seen in the pus from an abscess of the gum (*Boettcher*¹), in hypopyon (*Bizzozero*²), and in the contents of suppurating ovarian cysts (*R. v. Jakob*).

Some red blood-corpuscles are nearly always present in freshly secreted pus, and when blood in considerable quantity has been effused and afterwards its elements disintegrated, the discharge may be more or less amply tinged by admixture with blood pigment or hæmatoidin crystals.

Fatty particles and globules are seldom wanting, either free or combined with the protoplasm of the cells. The epithelium elements are comparatively few. In cancerous exudation from the pleural cavity vacuolated epithelial and fatty endothelial cells are commonly to be met with.

2. Fungi.—Modern research³ has established the fact that the formation of pus in animal organisms is effected almost entirely through the agency of micro-organisms, and that such bodies can nearly always be detected in the discharge with the aid of the staining methods which the recent advances of science have placed within our reach, where the microscope alone fails to disclose their presence.

Important experiments upon animals⁴ have shown also that suppuration may be caused by certain chemical substances, as cadaverin, croton oil, &c., independently of micro-organisms; and it is possible that the human system may be similarly affected by them. The products of bacterial life, toxines, and phytalbunins appear also to play an important part in the process of suppuration.

1. **Micrococci.**—Micrococci of varying form and size are very often to be seen in fresh pus.⁶ Fig. 144, which represents the appearance of a specimen taken from a pleural exudation and stained by Gram's method, affords an appropriate illustration. They are usually arranged in chains (Streptococci), occasionally in pairs (Diplococci).

Passet,⁷ proceeding on Koch's method, has cultivated no less than eight different forms of fungi from pus. When suppuration has continued for a long time in a cavity excluded from the air, micro-organisms are sometimes wanting in the pus. *Brieger*⁸ has observed *Staphylococcus pyogenes aureus* and *Streptococcus pyogenes* in pus from a woman with puerperal fever.⁹ The presence of micro-organisms hitherto mentioned is evidence of suppuration in the course of septic processes



FIG. 144.—Cocci from an Empyema, prepared by Gram's method (eye-piece III., objective Zeiss oil immersion $\frac{1}{4}$, Abbe's mirror and open condenser).

(comp. p. 54). *Bujwid*¹⁰ lately made the interesting discovery that the effect of grape-sugar upon the tissues is to promote the development of *Staphylococcus aureus* by diminishing their resistance, and so to favour the formation of pus. The tendency of diabetics to undergo suppurative processes, so long a matter of clinical observation, is explained in this way.

A blue colour has occasionally been observed on the surface of suppurating wounds. This is produced by colonies of *Bacillus pyocyanogenus* or of a fungus resembling it. The colouring matter has been isolated from such pus in combination with hydrochloric acid.¹¹

The detection of pathogenic fungi in pus is a point of great importance.

2. Tubercl-Bacillus.—The tubercle-bacillus has often been discovered in tubercular pus,¹² but the author has sometimes failed to find it even in the fresh discharge. Its presence, of course, is conclusive as to the nature of the disease, but its absence does not imply that no tubercle is present. It would appear that under certain conditions the bacillus rapidly disappears from fresh discharges (*Metschnikoff*).¹³

3. Bacillus of Syphilis. The bacillus discovered by *Lustgarten*¹⁴ in the pus of syphilis affords a valuable indication of this disease, but caution must be observed in identifying it, since *Alvarez* and *Tarel*¹⁵ have shown that certain secretions, as the smegma præputiale and vulvare, are apt to contain forms which closely resemble the venereal microbe. Such forms are to be discriminated by the behaviour of

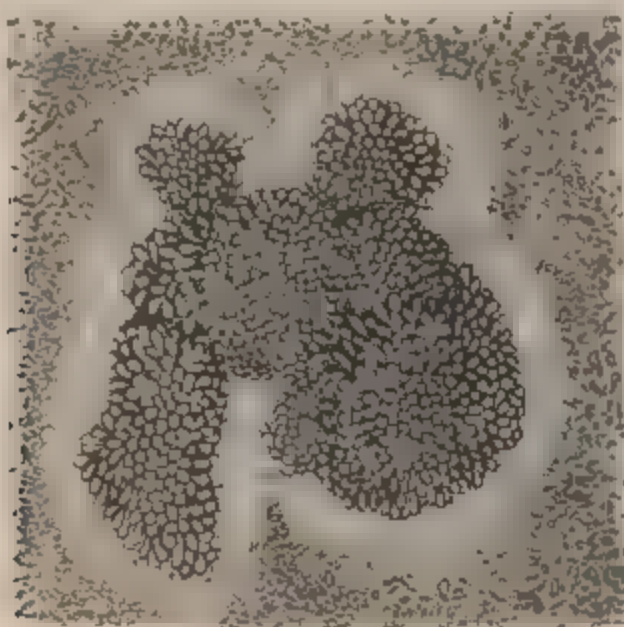


FIG. 145.—Actinomyces Granules in Glycerine from Actinomycosis of Pleural Cavity
(eye-piece II., objective IV., Hartnack).

stained preparations in presence of alcohol. In the case of the bacillus of syphilis, these are with difficulty and very slowly bleached by alcohol, whilst the microbe of smegma readily loses its colour under the action of that substance.

An important addition to our knowledge of the bacillus of syphilis was recently made by *Kamen*,¹⁶ who found the micro-organism in the sputum of a child of nine years. The character of *Lustgarten's* bacillus has been much questioned of late, and others have regarded certain cocci as the specific excitants of syphilis (*Kassowitz*, *Hochsteger*, *Disse*, *Taguchi*).

For the detection of the bacillus of syphilis *Lustgarten*¹⁴ proceeds as follows:—The cover-glass preparation is immersed in an Ehrlich-Weigert gentian violet fluid for 12-24 hours at the ordinary temperature. It

is then removed, rinsed for some minutes with absolute alcohol, and placed for ten seconds in a $1\frac{1}{2}$ per cent. solution of permanganate of potash, after which it is treated with a watery solution of pure sulphurous acid, and finally washed with water. Should it happen after this that the preparation still shows colour, it is again placed in permanganate of potash for three or four seconds, and afterwards in sulphurous acid until all colour has disappeared, the remainder of the process also being repeated. It is to be noted that other microbes, both pathogenic and innocuous, are stained by Lustgarten's process as well as that of syphilis.

*De Giacomi*¹⁷ has suggested a method which is very serviceable for the detection of the bacillus of syphilis. In this the dried cover-glass preparation is warmed for some minutes in an aniline-water fuchsin fluid, then washed with water to which a few drops of perchloride of

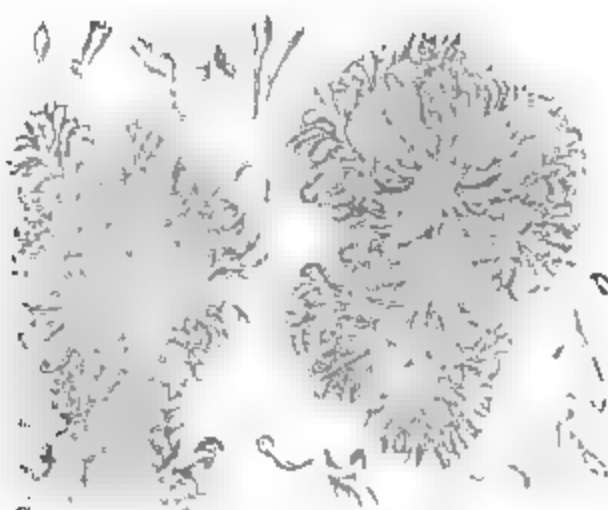


FIG. 146.—Actinomyces from a Case of Actinomycosis of Pleural Cavity (eye-piece III., objective, oil immersion $\frac{1}{2}$, Reckert; compressed).

iron solution have been added, and finally decolorised in a concentrated solution of that salt. The bacillus then remains of a red colour, while all other micro-organisms are bleached.

4. Actinomyces.—This parasite was first discovered by *Bollinger*¹⁸ in cattle, and afterwards in man by *Ponfick*¹⁹ and *Israel*.²⁰ In the former it gives rise to tumours of considerable size, but in man its proliferation is usually associated with chronic inflammation and the production of pus.

It would appear from numerous communications that have been made in recent years that actinomycosis is a disease of very wide distribution, and that amongst its symptoms is a severe form of angina, till lately obscure, to which the name of angina Ludovici has been given.²¹

[It usually affects the intestinal²² tract, but primary actinomycosis of the apices of the lung has been met with,²³ and six cases²⁴ have been recorded in which the disease had its seat in the brain. One of those by *Orlow*²⁵ is of

exceptional interest. *Anderson*²⁶ of Nottingham has lately recorded a case of actinomycosis of the face and neck, which was successfully treated by operation. [The best remedy, at present, for actinomycosis is sodium iodide.]

The pus of this disease is thin, viscous, and somewhat tenacious, and discloses to the naked eye small nodules of a grey or yellow colour, and about the size of a poppy-seed. Under a low power of the microscope these particles appear as a dense bunch-like aggregation of spherules, which under a higher magnifying power are seen to consist of pear-shaped, and radially arranged masses, which have a considerable refractive power (figs. 146, 147). Towards the centre of such masses the individual elements diminish in size and fade into a fine network of ramifying fibres. If one of the nodules be bruised, it shows in the first place numerous detached club-shaped forms, having a radiating disposition toward the periphery (fig. 146), and passing gradually at the centre

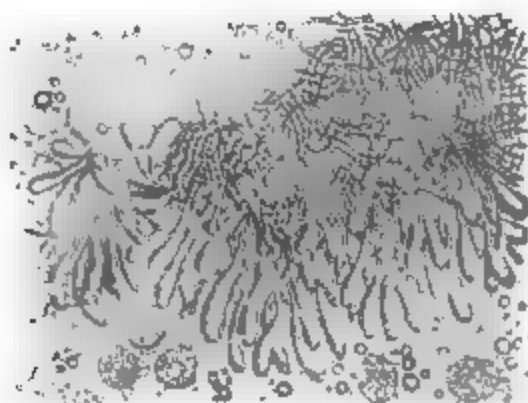


FIG. 147.—Actinomyces from Peritoneum (eye-piece III., objective, oil immersion, $\frac{1}{2}$, *Reichert*, unstained preparation).

into a sort of detritus; and in addition to these are a variety of other objects, club-shaped and of indeterminate appearance (degeneration types of the fungus), lying apart.

There was for a long time much doubt as to the botanical character of *Actinomyces*. Recently, however, it has been established²⁷ that it belongs to the genus *Cladothrix* and therefore to the class of fission-fungi. The characteristic club-shaped bodies must be regarded as degeneration forms of the fungus. In unstained microscopical preparations they may be seen sometimes to fade into the central fine network, and even occasionally to be enclosed in its meshes (fig. 147). [Crookshank²⁸ believes that *Actinomyces* is a fungus intermediate between the fission- and the higher fungi; and W. Hill²⁹ refers it to the Basidiomycetes.]

With the aid of staining methods (*Gram's* is the best) the individual threads of the network are readily distinguishable. They present a jagged or undulating contour, and result from the cohesion of a series

of minute spherical organisms, connected together by a remarkably delicate envelope. The centre, to which all the constituent threads of a group converge, is occupied only by a very dense network of this formation (fig. 148).

The pear-shaped bodies above alluded to may be more clearly defined by staining with *Weigert's*³⁰ process. For this purpose a solution is made by adding together 20 cc. absolute alcohol, 5 cc. concentrated acetic acid, and 40 cc. of distilled water, and to the mixture so much of the so called French extract of litmus as will give the fluid a dark-red colour, remaining ruby-red after repeated filtering (*Wedd's*³¹ litmus solution). In this solution the cover glass preparations are allowed to remain for an hour or so, then lightly rinsed with alcohol, and placed for two or three minutes in a 2 per cent. gentian-violet fluid, which should be boiled before use, and filtered after cooling.

If the specimen be now examined, it will be found that whilst the central mass of *Actinomyces* is colourless, the fungus threads are



FIG. 148.—Preparation from same case as last, stained by *Gram's* method (eye-piece IV., objective, oil immersion $\frac{1}{2}$, *Zeiss* Abbe mirror and open condenser).

stained a ruby-red. *Baranski*³² advocates staining with picro-carmine. In most cases, however, the character of this fungus can be determined by a simple microscopical examination. The physical qualities of the pus, and the discovery in it of groups of *Actinomyces* or the club-shaped degeneration forms of the parasite, are points which will aid the diagnosis. In particular cases it may be necessary to resort to Gram's staining method, and to observe the minute structure of the ultimate threads of the fungus, as noticed above (comp. p. 409). *Bugicid*³³ has obtained pure cultivations of *Actinomyces* by means of *Buchner's*³⁴ method (p. 448). Seen with the naked eye, these bear a close resemblance to cultivations of tubercle-bacillus.

5. Bacillus of Glanders. The specific micro-organism may be found in the pus from the ulcerated nasal passages in glanders. The method described in connection with the examination of the blood (p. 60) will serve for its recognition here.

Another method has recently been suggested (*Löffler*).³⁵ An aniline water-gentian-violet fluid or a concentrated alcoholic solution of methylene-blue may be employed, and immediately before use it is added to its own bulk of a (1 : 10,000) solution of potash. In the fluid which results the cover-glass preparation is immersed for five minutes. It is then removed and placed for a second in a 1 per cent. solution of acetic acid, tinged slightly yellow with oo-tropæolin. By means of another mixture, containing two drops of concentrated sulphurous acid and one of a 5 per cent. solution of oxalic acid in 10 cc. of water, the alkaline staining fluid may be decomposed and the preparation freed from colour. In this way the micro-organisms are very beautifully stained.

The bacillus of glanders may also infest the pus from an abscess. In any case, its character may usually be sufficiently determined in stained specimens by means of the microscope. When necessary, however, all doubt may be removed by observing the result of inoculating animals with the microbe. In the intermediate process of cultivation certain characteristic properties are disclosed. When the various other micro-organisms which are nearly always present in the pus have been eliminated by Koch's method (p. 448) the pure plate-cultivation crop made in nutrient agar-agar at 37° C. has the appearance of a greyish-white drop. The pure cultivation-product, when inoculated upon animals, as the mouse or guinea-pig, induces glanders. When engrafted on the potato and kept at a temperature of 35° C. the bacillus forms, in two or three days, a thin greasy coating of a brown colour. In coagulated blood-serum at a low temperature it develops after two or three days in the form of small scattered transparent drops which have a colour very much the same as that of the serum. The fungus is readily cultivated in glycerine agar-agar³⁶ and in nutrient milk-peptone.³⁷ When allowed to rest for some time longer, the cultivations are said to develop spores, but this point is not established (*Baumgarten*).³⁸

6. Bacillus of Anthrax.—Opportunity occasionally offers of examining the pus derived from a carbuncle in anthrax. The specific micro-organism is that described at p. 48. A certain familiarity with its life-habits will often facilitate its recognition, especially where it can be obtained only in very small numbers. The process by which these have been studied is in general the same as that employed for other pathogenic microbes. Thus the elimination of other forms is effected by Koch's plate (nutrient agar-agar and nutrient gelatine) and inoculation methods (see p. 448). When cultivated in a nutrient gelatine medium, the bacillus of anthrax develops in twenty-four to thirty-six hours, and forms minute points which are scarcely visible to the naked eye. With a lens the colonies are seen to be dark figures with an irregular undulating outline. Their peculiar shape becomes

more evident after the lapse of forty-eight hours, and later the cultivation liquefies, and from the dark central point it stretches in wavy strings over the entire surface of the plate.

The fungus does not liquefy agar-agar. When cultivated on sterilised potato, it forms a whitish-grey slimy patch of uneven surface, and extending but a few millimetres from the site of inoculation. Blood-serum cultivations form a superficial coating of white colour. Nutrient gelatine test-tube cultivations take the form of delicate white threads much interwoven, and gradually liquefy the gelatine; drop-cultivations of anthrax-bacillus in nutrient broth take the shape of long threads, in which, after a time, lustrous spots (spores) develop at regular intervals (comp. p. 49).

If the bacillus be imparted by inoculation to such animals as the mouse or guinea-pig, the animals exhibit the symptoms of splenic fever, and the micro-organism may be obtained from their blood.³⁹

7. Bacillus of Leprosy.—The nodes which are apt to form on various parts of the skin and mucous membrane in leprosy occasionally break down and ulcerate, with the formation of an abundant thin pus. In this, as indeed in the growths of leprosy generally, large numbers of bacilli may be found (*A. Hansen, Neisser*).⁴⁰ They have the form of minute rods $4\text{--}6 \mu$ long and 1μ in breadth, and have a close resemblance to the bacillus of tubercle (see p. 128). Like the latter, they stain in alkaline fluids, and are not bleached by subsequent exposure to acids. They are distinguished by the comparative readiness with which they stain, and by more easily taking up the colouring-matter from a simple watery solution of aniline dye.⁴¹ For their detection in the pus a suitable dry cover-glass preparation must be made (see p. 127), stained with Ziehl-Neelsen's carbol-fuchsin fluid, and again decolorised in acid (best with nitric acid) alcohol. [*Ehrlich's, Weigert's, and Gibbes' methods also serve.*]

Melcher and *Ortmann*⁴² claim to have produced leprosy in animals (rabbits) by inoculation with diseased tissue; and *Bordoni-Uffreduzzi*⁴³ to have cultivated the fungus from the spinal cord of lepers by inoculation in glycerine peptone-serum, glycerine blood-serum, and glycerine agar-agar. According to the latter, when cultivated by surface inoculation of nutrient substances, it forms ribbon-like cultivations with irregular borders, and of a light-yellow colour. On glycerine agar-agar it develops in small round whitish-grey cultivations, somewhat elevated at the centre, thinned at the edges and having a jagged outline.

[These statements need further confirmation. The attempt to cultivate the bacillus of leprosy has been made without decided success by many pathologists both in England and abroad. *Patrick Manson*⁴⁴ was one of the earliest to investigate the subject, and its literature

includes contributions by *Campana*,⁴⁵ *Kanthack* and *Barclay*,⁴⁶ *Brown Rake* and *Buckmaster*,⁴⁷ *Stalino*,⁴⁸ and *Byron*.⁴⁹ Dr. P. S. Abraham, the Secretary of the English Leprosy Commission, to whom Dr. Cugney is indebted for a personal communication, has examined specimens of the cultivations obtained by some of these observers, and he is of opinion that they are growths of a micro-organism of the skin which has nothing to do with leprosy.]

8. Bacillus of Tetanus. In view of the fact that the bacillus is found in the pus from wounds in cases of tetanus, and that much importance attaches to its recognition, some description of it is required here.⁵⁰ It occurs as delicate slender rods which have often a terminal spore, and thus present the appearance of bristles (see fig. 149). They

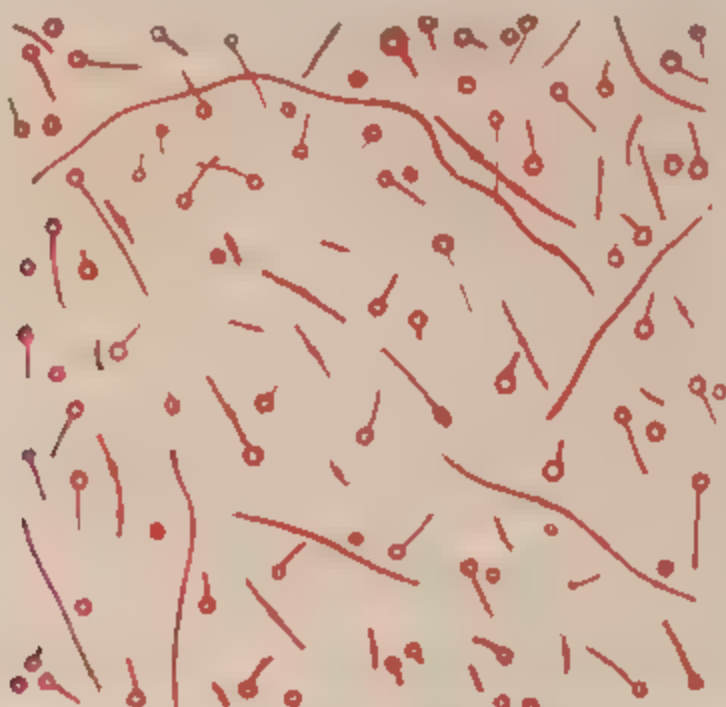


FIG. 149.—Bacilli of *Tetanus* (pure cultivation). Compensation eye-piece VIII, objective of immersion of Zeiss (from Koch).

stain with all the aniline dyes, and also with Gram's fluid. To obtain pure cultivations from pus, the cultivations first derived, which contain fungi of different kinds, are heated to 80° C. in a water-bath for from a half to one hour during a few days, after which they are inoculated on gelatine plates to which 1.5 to 2 per cent. of grape-sugar has been added, and these are kept in a hydrogen atmosphere (*Katalysat*) at 20-25° C. The resulting colonies are somewhat dense at the centre, with a more delicate and uniform periphery radiating from it. The preparation becomes fluid in time, and gas is evolved.

If a pure cultivation be inoculated under the surface of gelatine, growth begins near the surface of the gelatine, and the resulting cultivation is marked with faint radiating striae, or sometimes as thorn-like

processes projecting from it. Development is more rapid in agar-agar. After exposure of the bacilli for thirty hours to a temperature of 37° C. the spores already mentioned make their appearance.

A notice of the micro-organisms of pus would not be complete without taking account of the Cladothrix observed by *Eppinger*⁵² and others as occurring in abscesses. The author has recently detected Cladothrix bodies in pleuritic exudation, and inoculated them upon animals without result.

The number of micro-organisms capable of inducing suppuration is very great. It includes the typhus bacillus,⁵³ *Bacterium coli*,⁵⁴ *Gonococcus*,⁵⁵ and various cladothrices. Certain yeast fungi also, as *Busse*⁵⁶ has shown, may set up formidable and even fatal pyogenesis.

3. Protozoa.—Little is known as yet as to the presence of these parasites in pus. *Künstler* and *Pitres*⁵⁷ found numerous large spores with ten to twenty crescentic corpuscles in the pus from the pleural cavity of a man. These bodies closely resembled the coccidia which occur in the bodies of mice (comp. pp. 110, 216). *Litten*⁵⁸ observed cercomonads in the puncture-fluid, derived probably from the lungs. *Nasse*⁵⁹ has reported the presence of amœbæ in the pus from a hepatic abscess, and *Kartulis*⁶⁰ has observed them in the matter from submaxillary suppuration.

4. Vermes.—Filiariæ have been observed occasionally as occurring in the pus of tropical abscess of the liver.⁶¹ In temperate climates also abscesses result from an invasion of hydatids, and in such cases the pus contains entire echinococcus cysts, or fragments of the membrane and hooklets. *Babesiu*⁶² found filaria in the gastro-splenic omentum. They were probably the same as *Grassi's*⁶³ *Filaria inermis*. With these should be mentioned the bodies resembling filaria which *Sarcani*⁶⁴ reports as found by him in the suppurating parotid of a woman.

It may be mentioned here that v. *Bondzynski*⁶⁵ distinguishes the cholesterolin of human faeces from other forms and gives it the name of coprosterin.

5. Crystals.

1. Cholesterin.—Crystals of cholesterin are very rarely to be found in fresh pus—more commonly in that derived from cold abscesses, and most abundantly in foetid discharges and in suppurating ovarian cysts. For their character and recognition the reader is referred to pp. 136, 240, 289.

2. Hæmatoidin.—This body exhibits the same variety of form here as in the sputum, the urine, and the faeces (pp. 135, 233, 280). Its presence invariably points to a previous haemorrhage. It is specially abundant in the case of suppurating hydatid cysts.⁶⁶

3. Fatty Needles.—These are of very many shapes and forms,

occurring sometimes singly, and sometimes in groups and clusters. They are the outcome of a degenerative process, and show that the pus in which they are found has been long formed. Very perfect margarin needles may be obtained from gangrenous pus (fig. 150).

4. Triple Phosphate.—Crystals of triple phosphate are a common constituent of the pus (see p. 281). Crystals of carbonate and of phosphate of lime are also met with, and most commonly in fœtid pus.

III. CHEMICAL EXAMINATION OF THE PUS.—It rarely happens that much aid accrues to diagnosis from the chemical examination of the purulent discharges. The albuminous substances which may commonly be recognised are serum-albumin, globulin, and especially peptone (*Hofmeister*⁶⁷) in large quantity. Fat also is found.⁶⁸ The peptone is derived from the cells, not from serum. For the means of detecting these bodies the reader is referred to p. 296.

Fresh pus always contains glycogen, and traces of grape-sugar are seldom wanting. In testing for the latter, the pus should be boiled with an equal weight of sodium sulphate, to free it from albumin, filtered, and the filtrate treated in the manner previously described (p. 317).

In cases of jaundice the pus may contain bile-pigments and biliary acids.

Considerable quantities of nuclein, fats, cholesterin, and a number of inorganic salts, notably phosphate and chloride of sodium (*Miescher*, *Naunyn*⁶⁹), are constantly to be found.

In three specimens of pleuritic exudation which he has investigated the author found abundance of acetone, and it would appear from a private communication of *Professors Baumann* and *Baumler* that that body is commonly a constituent of pus. In another specimen submitted to him by *Dr. R. Paltauf*, whose attention had been attracted by the evident odour of acetone, it was found by the author to be plentifully present; and the researches of others have shown that acetone is frequently present in large proportion in exudation fluid. Fœtid pleural exudation often contains sulphuretted hydrogen. The method for its detection is described at p. 381. There, as in certain forms of hydrothionuria, fungi possessing the property of liberating sulphuretted hydrogen may be isolated from the fluid.

*Guttmann*⁷⁰ found indigo-producing substances in exudations. The author has repeatedly established the presence of fatty acids,—acetic, formic, and butyric acids. Pus contains further a trace of uric acid and several xanthin bases. Much interest attaches to the occasional presence of guanin (*R. v. Jaksch*).⁷¹

2. SERO-PURULENT EXUDATIONS.

Sero-purulent closely resemble the purulent discharges in their chemical, physical, and morphological character. They are distinguished chiefly by the relatively smaller proportion of extractives which they yield. They invariably point to antecedent inflammation.

3. PUTRID EXUDATIONS.

Discharges of this character are brown or brownish-green in colour, and have a penetrating and disagreeable odour. Their reaction is usually alkaline. Microscopically they exhibit much-shrunken leucocytes and an abundance of crystals, chiefly of fat, and of cholesterin and haematoxin in lesser proportion. They also contain a profusion of fission-fungi of different kinds (see fig. 144).



FIG. 150.—Pus from Putrid Empyema (eye-piece III., objective 8A, Reichert).

4. HæMORRHAGIC EXUDATIONS.

A hæmorrhagic exudation contains an abundance of red blood-corpuscles, and often also a considerable quantity of dissolved hæmoglobin. Endothelial cells loaded with fat are almost invariably to be found, and when those occurring in pleuritic fluid exhibit the glycogen reaction in a marked degree, they favour the assumption of carcinoma (*Quincke*).⁷² A positive diagnosis upon this point results from the further discovery of cancer-cells.

In the absence of determinate (specific) elements, as cancer-cells, tubercle-bacillus, &c., no very definite conclusion can be drawn from the hæmorrhagic character of the discharge, since many different processes may be associated with the effusion of blood. Still in the case of pleuritic fluid, where scurvy and cancer of the pleura can be excluded, the appearances would point to tuberculosis.

5. SEROUS EXUDATIONS.

Fluids of this class are almost quite clear and more or less deeply tinged yellow; they coagulate on standing (for twenty-four hours), and yield a clot which is usually rich in fibrin. Microscopically, they exhibit scattered red blood-corpuscles in usually fairly good preservation, but sometimes much attenuated, a number of leucocytes, some fatty globules and endothelial cells, separately or in groups. In addition to these are sometimes cells ranging from $7-30\ \mu$ in diameter, and formed of very small droplets. These may contain two or three large cavities (*Bizzozero*).⁷³

There is reason to believe that micro-organisms very frequently infest the serous exudations, but the subject calls for further elucidation. It would appear, at any rate, that fungi occur oftener in them than in the transudations which so closely resemble them in their physical and chemical properties. The fluid obtained from the pleural cavity in tubercle of the pleura with breaking down of tissue may contain the specific bacillus of that disease. In cases, however, where no discharge of tubercular matter has taken place into the pleural cavity, the tubercle-bacillus will not be found.

Cholesterin crystals occur in serous discharges of old standing. Chemically, serous exudation fluid contains serum-albumin and globulin, but no peptone. Sugar in small quantity is invariably present,⁷⁴ and acetone from time to time. A notable fact is the frequent or perhaps invariable occurrence of uric acid in exudations of this kind.

The density of the fluid is clinically a fact of much consequence. It may be estimated by means of a pycnometer, or with an accurate aërometer, regard being had at the same time to conditions of temperature. The sp. gr. will usually be found to exceed 1.018 (*Reuss*).⁷⁵

6. CHYLOUS EXUDATIONS.

Peritoneal exudation is commonly characterised by the abundance of fatty matter which it contains. This also is especially true of discharges depending upon obstruction of the thoracic duct.

The appearance of chyle, however, is sometimes misleading, since it has been shown that this character belongs in general to pathological fluids of low sp. gr., especially to such as owe their origin to passive congestion (*F. A. Hoffmann*).⁷⁶

*Boulengier*⁷⁷ draws a distinction between chylous and chyliform exudation, limiting the first of these terms to the case in which chyle is actually discharged into the peritoneal cavity, while the second implies only that the fluid has the properties of chyle. Chylous effusion

is very rich in fat. A specimen of chylous pericardial fluid was found by *Hasebroek*⁷⁸ to contain 10 per cent. of the latter.

For the discrimination of the different forms of pleuritic effusion, the bacteriological evidence is of much importance.⁷⁹ Thus the absence of micro-organisms points to a tubercular origin when the exudation is purulent; sero-fibrinous exudations are also usually free from fungi, and cases of empyema occur in which only *Staphylococcus pyogenes*, and still more often *Streptococcus pyogenes* (*Ludwig Ferdinand of Bavaria*) are found. Exudations occurring in the course of pneumonia often exhibit Fränkel's pneumonia-coccus. The bacteriological investigation of pus in other organs also is a matter of great interest, as is proved by the researches of *Wertheim* in *Schauta's* clinic.

In general it is a difficult task to determine whether a particular fluid effused into one of the cavities of the body is due to inflammation (exudation) or to simple obstruction (transudation). In such cases a probable conclusion may be based upon the sp. gr. of the fluid (*Méhu, A. Reuss*).⁸¹ A further point of interest in this connection is the fact that inflammatory fluids are relatively rich in fibrin (for the estimation of this body see p. 313) and in dry residue products.⁸² A large percentage of proteids points the same way. For the estimation of these, *Kjeldahl's* process may be adopted.⁸³ It must be borne in mind that neither the sp. gr. nor an abundance of proteids is absolutely distinctive of an inflammatory origin (*A. Ott*),⁸⁴ and these must be taken in conjunction with other particulars to form a just conclusion. The author in collaboration with *Ott* has ascertained that pus holds about 8 per cent. albumin; transudations contain less, 4-5 per cent.; exudations more than this, 6-8 per cent.

B.—TRANSUDATIONS.

Transudation fluids may be serous, sanguous, or, in rare instances, chylous. Their sp. gr. is in general lower than that of inflammatory effusion into the same cavity. They are always alkaline,⁸⁵ and for the most part of a yellow colour.

Microscopically they exhibit but few tissue elements—fewer than, but in other respects similar to, those of serous exudation. It should be noted that in serous pleuritic effusion a large quantity of endothelium is often detached. This must not be taken to imply endothelial proliferation in a new growth (carcinoma), unless blood be also present, when such an inference acquires additional support.⁸⁶ Chemically, transudation products exhibit much albumin, and generally sugar.⁸⁷ For the detection of the latter the process detailed at p. 319 may be

employed. They are always free from peptone. *Paijkull*⁸⁸ states that nucleo-albumin is absent from non-inflammatory transudations.

Such fluids are distinguished from exudations chiefly by their low sp. gr. and the difficulty with which they coagulate. The distinction, however, is not easy to make in many instances.⁸⁹

The author would observe that in six specimens of transudation fluid and serous exudation, which were entirely free from blood-corpuscles and dissolved blood-pigment, he succeeded in separating no small quantity of urobilin. This body is commonly to be found in transudations and exudations, and uric acid is always present in them. In the transudation from a case of cirrhosis of the liver *Moscatelli* recently found allantoin.⁹⁰ For the detection of uric acid the process described at p. 85 may be applied. The occurrence of urobilin in exudations and transudations has been established by *Ajello*.

C.—CONTENTS OF CYSTS.

The physician has often to decide whether a particular specimen of fluid withdrawn from the body by aspiration or puncture is inflammatory in its origin or due to passive congestion, and finally whether it is derived from a cyst. Such a question is least apt to arise in exploration of the pleural cavity, but in connection with abdominal symptoms it is frequently very urgent to be able to give a definite answer. And this is not always easy; at times, indeed, it is impossible.

The cysts with which we have to do in this way are hydatid and ovarian cysts, and, in rare instances, cystic kidney and pancreatic cysts.

1. Hydatid Cysts.—The fluid obtained by puncture from a hydatid cyst is clear, alkaline, and usually of low sp. gr., 1.006–1.010. It contains a small quantity of a reducing substance (grape-sugar), very little albumin, and abundance of inorganic salts, as sodium chloride.⁹² Succinic acid and inosite have also been detected. The microscopical appearances are very characteristic. Amongst these are the hooklets (see also p. 134) and portions of echinococcus membrane, so readily distinguished by its transverse striation and uniformly granular inner surface (p. 134, fig. 63). Scolices may also be seen, and are known by the two circles of hooklets and four suctorial discs on the anterior aspect (head) and the sack-like hinder part, which is separated from the head by an annular constriction. If the cysts have suppurated or be filled with blood, as sometimes happens, chemical examination will usually throw but little light upon their character. An absolute diagnosis is possible only when hooklets or shreds of membrane have been seen in the fluid. Consequently it is well to receive the latter in a conical glass, and to carefully examine the sediment for these bodies.

Hydatid cysts often contain haematoxin crystals (see p. 233).

2. Ovarian Cysts.—The fluid obtained from ovarian cysts is remarkably variable in its character.

It is generally to be distinguished from inflammatory and congestion fluids by its high sp. gr., which ranges from 1.020 to 1.026. Its reaction is alkaline, and it has little tendency to coagulate.

It is further marked by the great abundance of tissue débris which it contains, and, from the nature of the cells which preponderate, information may be drawn as to the kind of cyst from which it has been taken.

Instances, however, occur in which the fluid from an ovarian cyst shows nothing by which it may be known from that of ascites, and exceptionally, it has a sp. gr. even lower than that of a transudation fluid.

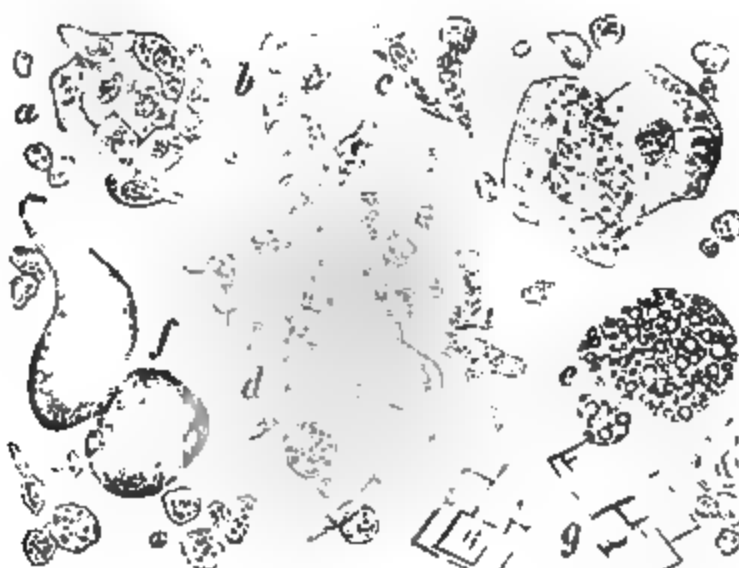


FIG. 151.—Contents of an Ovarian Cyst (eye-piece III., objective 5A, *Reichert*).
a. Squamous epithelium cells; *b.* Ciliated epithelium cells; *c.* Columnar epithelium cells.
d. Various forms of epithelial cells; *e.* Fatty squamous epithelium cells; *f.* Colloid bodies; *g.* Cholesterol crystals.

According to *Schatz, Gruerow, and Westphalen*,¹⁸ a low sp. gr. with little albumin points to a cyst of the broad ligament.

When haemorrhage has taken place into the cyst, its contents may vary in colour from red to a chocolate-brown, and be very turbid. The microscopical examination of ovarian fluid shows a very variable quantity of red and white blood-corpuscles, and many forms of epithelium, squamous, columnar, and ciliated (fig. 151, *a*, *b*, *c*). These cells, however, are rarely well preserved, but are for the most part far advanced in fatty degeneration, and often with difficulty recognisable. Colloid concretions (fig. 151, *f*), in all probability derived from epithelium, are invariably found in the so-called colloid cysts.

Certain forms of ovarian cysts may be readily differentiated by a microscopical examination of their contents. Thus in dermoid cysts

may be seen, besides squamous epithelium, hairs, and crystals—cholesterol, fatty, and haematoxin. Chemical investigation, too, throws much light upon the character of an ovarian fluid. It will be found as a rule to contain albumin, and always metalbumin (paralbumin), and it is this body principally which renders it turbid and stringy (*Hammarsten*).⁹⁴

To Test for Metalbumin.—The fluid is mixed with three times its bulk of alcohol and allowed to stand for twenty-four hours, when it is filtered, the precipitate squeezed out, and suspended in water, which is again filtered. The filtrate is opalescent, and has the following characters:—(a.) On boiling it becomes turbid, but does not form a precipitate. (b.) No precipitate forms with acetic acid. (c.) Acetic acid and ferro-cyanide of potassium render the fluid thick and impart to it a yellow tint. (d.) Millon's reagent, on boiling, yields a bluish red colour. (e.) Concentrated sulphuric and glacial acetic acid yield a violet colour (*Adamkiewicz*). (f.) *Huppert*⁹⁵ has pointed out that when the fluid containing metalbumin is boiled with sulphuric acid it yields reducing substances, and this he regards as one of its most characteristic properties. It must be noted that metalbumin is a constituent of other pathological fluids besides that of an ovarian cyst. These cysts, and especially dermoid cysts, also hold a considerable quantity of cholesterol in solution. Cystic fluids have been frequently shown to contain diastatic ferment in appreciable quantity.⁹⁶

3. Cystic Kidney.—In all cases where a sufficient specimen can be obtained, the fluid from a cystic kidney (hydronephrosis) may be at once identified on the ground of its chemical properties and microscopical appearances. An important point is the presence of renal epithelium, and this should be carefully sought for. Then the determination of urea and uric acid in large quantity indicates a connection with the kidney; but it must not be forgotten that these bodies occur more or less plentifully in ovarian cysts, or may be supplied to them when they communicate with the urinary tract by an abnormal channel.

It has been said that for the purpose of diagnosis the greatest significance attaches to the presence of epithelium from the urinary tubules; and inasmuch as this usually occurs very sparingly in the fluid from a cyst, the latter after removal by puncture should be allowed to settle and the sediment carefully examined. Such of the clinical symptoms of cystic kidney as call for notice here are very various. *P. Wagner*⁹⁷ observed that the urine is often scanty. Albuminuria and intermittent haematuria, symptoms of sufficiently vague import, also occur.

4. Pancreatic Cysts.—The fluid from a pancreatic cyst is of low sp gr.—1.010–1.012 (*Karewski*⁹⁸), 1.022 (*Hofmeister*⁹⁹) 1.028 (r.

*Jaksch*¹⁰⁰)—and usually, though not always, has an admixture of blood. In the author's experience the blood pigment is in the form of methæmoglobin. There is also present abundance of cholesterol. Of proteids there is serum-albumin; rarely mucin is found. Metalbumin is absent. The fluid also contains diastatic ferment, but the fact has by itself no weight in diagnosis, since that body is very frequently met with elsewhere (see p. 244). It is only when the sugar resulting from its action is found to be maltose that any significance attaches to its presence (see p. 334).

Such a fluid has the property of digesting albumin without the addition of acid, and it is this property which lends itself chiefly to the purpose of diagnosis. To test it, according to *Boas*,¹⁰¹ the fluid may be added to milk, and after precipitation of the casein the biuret test is applied. If the reaction is obtained, the fluid is shown to have the peptonising property, and it may be inferred that it is derived from the pancreas, since no other cystic fluid is known to dissolve albumin in alkaline solution. Of less moment is the property of such fluids to emulsify fat and to disengage carbon dioxide gas on the addition of acids. The importance of the characters here described is in most cases greatly impaired by the fact that the larger and the older the cyst, the less do its contents exhibit the physiological peculiarities of the pancreatic juice (*Wölfler*).¹⁰² It follows that where the clinical symptoms point to a cyst of the pancreas, it is not admissible to reject the inference on the ground that the fluid is devoid of the tryptic character, and to this extent the diagnostic value of these chemico-physiological observations is necessarily curtailed.

D.—THE SECRETIONS FROM FISTULA.

Where these consist of a purulent, or simply serous fluid, discharging through an unnatural outlet, their diagnosis may be based upon the considerations already dealt with under the headings Transudations and Exudations. A greater, or at least, at present, a more physiological interest attaches to the appearances in those cases where fluids are discharged by an unnatural opening, which might be assumed to communicate with the intestine, because the fluid closely resembles the intestinal fluid in its physiological properties, while fuller investigation shows that it is derived from cavities lined by secreting glandular epithelium.

The author has analysed the secretion from a case of this kind which *Professor Wölfler* observed and operated upon. The fluid had an acid reaction, and contained albumose and peptone in considerable quantity, pepsin, and a ferment which changed maltose into grape-sugar; no diastatic ferment.¹⁰³

The maltose employed in this research gave Trommer's reaction only very faintly (see p. 318). When it acted upon the secretion at 40° C., both Trommer's and Nylander's tests gave positive results.

Small quantities of free hydrochloric acid were perhaps also present—the Congo-red and benzo-purpurin tests gave feeble results—but there was no sugar, urea, bile-pigment, or urobilin. Of inorganic salts there were chlorides.

From this statement it appears that the fluid in many of its characters resembled the mixed secretion of the intestinal tract.

CHAPTER IX

THE SECRETIONS OF THE GENITAL ORGANS

I. THE SEMINAL FLUID.

1. Naked-Eye Appearances of the Semen.—The semen is a thick, white, and somewhat opaque fluid, of slightly alkaline reaction. It is tolerably tenacious, and resists the pressure of the cover-glass. It owes this property to the presence in it of a gelatinous substance, which, under the microscope, appears to be hyaline, and encloses innumerable cavities of various sizes. The semen has a peculiar odour, which is derived, according to *Fürbringer*,¹ from the prostatic fluid, and depends upon the large proportion of compounds of Schreiner's base (æthylenimin) (see p. 135) which it contains.

2. Microscopical Examination of the Semen.—The spermatozoa, which are to be seen in great numbers in the semen, are thread-like bodies about $50\ \mu$ in length, and provided with a head $4.5\ \mu$ long, and compressed in one plane, so as to appear club-shaped when seen from the side. They exhibit very lively movements, but their motility is rapidly destroyed by the addition of water, drying, &c. They are present only in semen and in fluids with which the latter is mixed, and obviously they may possess a great interest in the diagnosis of certain morbid conditions. It may happen that the physician will have to examine the semen to settle a question of sterility. A persistent absence of spermatozoa (azoospermia) will show that the individual is incapable of procreation, and this may occur whilst the other signs of sexual power are retained. Of forty cases of sterile marriages, *Kehrer*² found that azoospermia was the cause in fourteen. It is very important to distinguish the persistent condition from the temporary absence of spermatozoa, which occurs as a result of excessive and repeated intercourse. Under such circumstances the fluid ejaculated consists almost entirely of prostatic secretion (*Fürbringer*).³

In addition to spermatozoa, the semen exhibits certain cells—large and small, finely granular testicle-cells, with one or more nuclei—some columnar and squamous epithelium, a few large round hyaline bodies, lecithin corpuscles, and stratified masses of amyloid substance, which

are finely granular within, and usually enclose a central kernel—these are derived from the prostatic secretion; and, finally, a few leucocytes—which usually have two nuclei—and spermatic crystals. Some red blood-corpuscles may also be seen.

Certain pathogenic micro-organisms, and especially tubercle-bacillus, may be present in the secretion from the genital tract. They are usually discharged with the urine. The seat of the disease may be determined in such cases by other signs, as swelling of the testicle and epididymis, &c., comp. p. 275).

In some pathological conditions the semen may be coloured a chocolate-brown from the presence of a quantity of amorphous blood-pigment. This happens especially in old people and in persons who have often suffered from orchitis.

A very particular interest attaches to the spermatic crystals. In their appearance and chemical properties they very closely resemble the



FIG. 152.—Microscopical appearance of the semen (human). eye-piece III, objective 8A, *Reichert*.
 a Spermatozoa; b Columnar epithelium cells; c Bodies enclosing lecithin granules; d Squamous epithelium cells from the urethra; e Testicle cells; f Amyloid corpuscles; g Spermatic crystals; h Hyaline globules.

crystals which have been already described as occurring in the blood, sputum, and faeces (see p. 134). *Furbringer* has shown that while the basic compound is derived from the prostatic fluid, the phosphoric acid combined with it is furnished by the other component of the semen—the spermatic or testicular secretion. The crystals' form immediately and in great abundance, when a 1 per cent. solution of ammonium phosphate ($(\text{NH}_4)_2\text{HPO}_4$) is added to the pure prostatic fluid, and their presence in large numbers, therefore, under all circumstances, indicates prostatorrhœa (see p. 424).

It follows, therefore, that such crystals are not characteristic of the semen, and the admixture of that secretion with some fluid or in a dried discharge can be established only on the discovery of spermatozoa. For this purpose, when dried, they must be dissolved out in water from the discharge containing them.

3. Chemical Examination of the Semen.—But little informa-

tion can be derived in this way. According to *Miescher*, the fundamental constituent of spermatozoa is nuclein. Globulin and serum-albumin have been found in the semen, and it is very rich in inorganic substances. *Posner*⁴ asserts that albumose is also present.

II. SECRECTIONS OF THE SEXUAL ORGANS OF THE FEMALE.

1. Mammary Secretion (the Milk).—During the entire period of gestation, and especially from the third month of pregnancy onwards, a thin, whitish, and more or less turbid fluid may be obtained by pressure from the breast. This is a fact of great importance, and the existence of such a secretion is by itself strong evidence of pregnancy.

The fluid in question, when examined microscopically, presents in the first place a great number of strongly refractive bodies of very irregular size. These are called colostrum-corpuscles; they are fatty

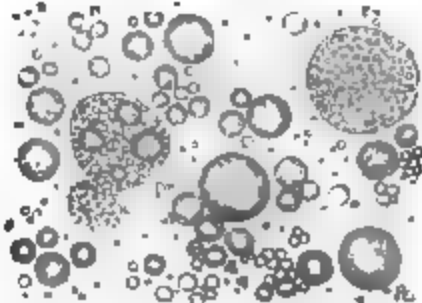


FIG. 153.—Colostrum of a Woman in Sixth Month of Pregnancy (eye-piece III., objective 8A, Reichert).

in character, and are usually aggregated in groups. In addition to these are to be seen a few leucocytes and some epithelial cells from the inner surface of the ducts.

Immediately after confinement the colostrum-corpuscles disappear from the milk, and eight or ten days after parturition are no longer to be found. *Czerny*⁵ supposes that colostrum-corpuscles are lymphoid cells whose function it is to take up and alter the unused milk-globules, which they then transfer from the acini of the gland to the lymphatics. In their place is seen a profusion of fatty globules of irregular dimensions, and together with these certain particles (*Hoppe-Seyler*) which consist of casein and nuclein.

In diseases of the breasts, and especially in cases of abscess and inflammation during suckling, the milk is apt to exhibit an intermixture of leucocytes.

Micro-organisms occur in the secretion in connection with certain morbid states. Thus *Escherich*⁶ found fungi in the milk of a woman suf-

ferring from septicæmia, and these on cultivation proved to be pathogenic. Pathogenic Staphylococci have also been isolated by Koch's cultivation-process from the milk of a woman with facial erysipelas (*Karlinski*).⁷ Similarly the author has detected micro-organisms, and especially coccii, which stained by Gram's method, in the milk of a patient with puerperal septicæmia.⁸ Further observations on the presence of fungi in human milk have been published by *M. Kohn* and *H. Neumann*.⁹

There can be no doubt that milk may contain the bacillus of tubercle, and indeed almost any of the pathogenic fungi. For the detection of tubercle bacillus the method of *Arnell Kunt*¹⁰ may be employed.

The milk of some of the lower animals has been known to be infested by non-pathogenic micro-organisms (*Bacillus cyanogenus* and *Micrococcus prodigiosus*), from which it may derive an abnormal blue or reddish tint (*Veelzen, Hueppe*).¹¹

The chemical constitution of the milk varies under different conditions both of health and disease. The milk of sick women is generally less rich in fat, and the proportion of lactose is diminished. Neither bile-pigments nor biliary acid have yet been satisfactorily demonstrated in the secretion of jaundice (*v. Jaksch*).¹² The albuminous constituents of human milk are serum-albumin, casein, and nuclein. It also contains milk-sugar and fats. For the detection and estimation of the latter, the method described for the same purpose in the chapter on urine may be employed. Special processes for the quantitative analysis of the milk may be found in *Hoppe-Seyler* and *Thierfelder's* work; *Kjeldahl's* process is applicable to the estimation of albumin.

The examination of the milk of wet-nurses is a point of practical interest for the physician. It should be tested carefully as to its naked-eye and microscopical characters, and in all cases chemically analysed. In addition to this, information may be derived from the employment of the bacteriological methods. And it is highly expedient that the milk, whether of healthy or diseased women, should be submitted, wherever possible, to Koch's plate-cultivation processes, so as to ascertain the absence of fungi.

2. Vaginal Secretion.—This, under ordinary circumstances, is a thin fluid with an acid reaction. It contains a few large leucocytes, each with a single nucleus, and squamous epithelium cells, which, for the most part, are covered with microbes. In vaginal catarrh the number of leucocytes is greatly increased, and some red blood-corpuscles may be visible.

In cases of ulcerating carcinoma implicating the vagina or the vaginal portion of the uterus a copious fetid discharge takes place, and in this the characteristic large cells of cancer may be found (fig. 154).

*Hausmann*¹⁴ detected fatty needles in the vaginal mucus.

Among the parasites which have been found in this situation are :—

1. Yeast and Fission-Fungi. — Various fungi belonging to these classes infest the vagina. Thrush-fungus vegetations also have been seen there. The vaginal secretion normally (*Winter*¹⁵) and during confinement (*Duxlerlein* and *Samschin*¹⁶) contains fission-fungi, as, e.g. *Staphylococcus pyogenes albus*, *aureus*, and *citreus*. *E. Bumm*,¹⁷ on the other hand, maintains that the secretion in health is free from pathogenic micro-organisms. In purulent catarrh (gonorrhœa) the micro-organisms which cause infection in the male are found, but not always in proportion to the virulence of the contagion, because the vaginal secretion possesses bactericidal properties. Finally, it is sometimes important to examine the secretion for tubercle-bacillus and gonococci by the prescribed methods.

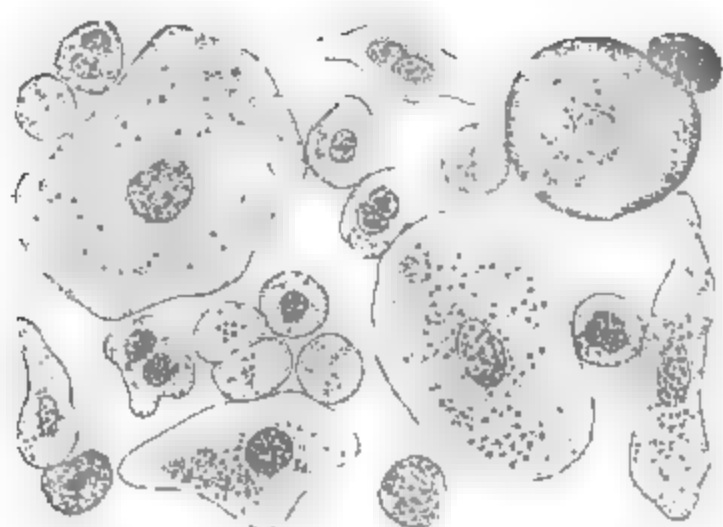


FIG. 154.—Preparation of Vaginal Secretion in a Case of Cancer of the Cervix Uteri
(eye-piece III., objective 8A, *Reichert*).

Of the chemical constitution of the vaginal mucus little is known. It has been said to contain trimethylamin (*Hilger*).¹⁸

2. Trichomonas vaginalis. — This parasite is an infusorium of oval shape, and about $10\ \mu$ in length. It has a long caudal appendage, three flagella, and a lateral row of cilia.

3. The Uterine Secretion.

1. Menstruation. — At the outset of menstruation there is an increased discharge of vaginal secretion. Later on there is mixed with this an abundance of red blood corpuscles and prismatic epithelial cells laden with fat from the interior of the uterus. The proportion of blood-cells begins to diminish after a few days, and leucocytes preponderate towards the close of the period. At this time also epithelium and a large quantity of fatty detritus are discharged.

2. The Lochia. — The fluid discharged during the first few days after parturition is thin and of a red colour. In addition to red and white

blood-corpuses it exhibits uterine and vaginal epithelium. Later, whilst the red corpuscles diminish in number, the leucocytes and epithelium increase, and the discharge assumes a grey or even white colour.¹⁹ Microbes are always plentifully present, even in the absence of septicaemia. According to *Döderlein*,²⁰ healthy lochia are free from fungi, while in disease he found them without exception to contain *Streptococcus pyogenes*. These statements are confirmed by *Thomen*.²¹

A point of special importance in diagnosis is the examination of the uterine secretion for the pathogenic fungi, to which reference has been already made. The secretion may be removed for the purpose by means of a tampon.

CHAPTER X

METHODS OF BACTERIOLOGICAL RESEARCH

THE great practical importance which this subject has acquired of late makes it necessary that the physician should be familiar with the processes for ascertaining the presence of micro-organisms.

In all cases where micro-organisms may be the exciting cause of disease, his first task will be to detect them in the secretions or natural fluids by means of the appropriate methods for staining them.

When this is done, it remains in many instances to discover the micro-organisms in particular situations, cells or tissues, so as to exclude a mere misleading coincidence.

Farther, the micro-organisms are to be cultivated outside the system ; so that, where their morphological character and their behaviour to staining substances are not sufficiently distinctive, the requisite inferences may be drawn from their mode of growth and development.

Finally, it is possible by experiments upon animals to settle definitely whether the transmission of a pure cultivation of the micro-organisms will produce symptoms more or less closely resembling those attributed to their agency in man.

The staining methods at our disposal and the perfection of our optical instruments in many cases render the detection of micro-organisms easy, but their cultivation and the transferring of them to animals is often very difficult. Thus in the case of many diseases, micro-organisms have been discovered under such circumstances as to leave no doubt that they were the exciting cause, while every effort towards their cultivation and transmission to animals has failed of success. This, however, has been achieved in the case of a number of pathogenic fungi, as, for instance, the bacilli of glanders, anthrax, tubercle, cholera, leprosy, tetanus, actinomycosis, and typhoid fever.

It is no longer necessary for diagnosis to pursue the inquiry in every instance through its entire course (detection, cultivation, and transmission to animals) ; but in some diseases, as tuberculosis, it is sufficient to note the characteristic effect of staining substances. In others, again (as relapsing fever and, occasionally, anthrax), a simple microscopical examination will serve, without the application of staining

processes. In doubtful cases of anthrax, the diagnosis may be settled securely by the direct transmission of the blood to animals.

With reference to Asiatic cholera, the mere detection of the fungus in the stools is never sufficient, but it must be isolated by Koch's cultivation methods, when it will be recognised by its mode of growth. With the progress of science we may learn to know a definite fungus as the cause of each of the infectious diseases; but even then, when all the conditions indicated above have been complied with, our labour will not have ended. It will remain to extend our acquaintance with the life-history of the parasites, so as to determine the sources of nitrogen and of carbon and the inorganic salts upon which they depend for their growth. It is only when this is done that a secure foundation will be laid for a system of rational therapeutics.¹

A short account will be given here of the methods employed in such researches. It will naturally begin with a description of the apparatus, and in the first place of the microscope.

I. THE MICROSCOPE.

The shape, size, and adjuncts of the body or stand of the microscope itself are in general of little moment. Habit will determine, in most cases, whether the tube used shall be worked by a screw or with the hand. For the examination of plate-cultivations, however, the former will be chosen. Neither is it essential that the stand should be jointed, *but it is absolutely necessary that it be faultless in its construction. It must also be capable of adjustment for use with the most powerful objectives, and with an Abbe's condenser or equivalent arrangement.*

The stage must be sufficiently large and firm, and the opening in it as large as possible, so that a plate-cultivation, for instance, may be inspected easily with a low power.

For bacteriological investigations, as has been already observed, an Abbe's or other condenser adjusted movably to the microscope-stand is needed. The principle of such an instrument is this: The rays of light reflected from the mirror of the microscope are passed through the principal axis of a lens interposed between the mirror and the objective in such a direction that they fall upon the object, which thus receives a cone of light as highly concentrated as possible. If narrow diaphragms be introduced between the mirror and the collecting lens, an illumination of the image is obtained similar to, but probably somewhat more intense than, that where narrow cylindrical diaphragms are used. The edges of the image are in all cases well defined, even in unstained preparations, and such a condenser may be employed with advantage for histological work. If the diaphragms be removed and

an open-condenser light consequently employed, the outlines disappear and become entirely undistinguishable (*Koch*).² Under these circumstances nothing else can be distinctly made out with uncoloured preparations. With stained preparations, however, it is far otherwise, and in this lies the valuable application of the open-condenser light as discovered by *Koch*. The outlines, in so far as they depend for their appearance on a distinction in the refractive properties of the object (corresponding to a structural difference amongst its parts) and those portions which are but lightly stained, are lost to view, while the deeply-stained particles, as the coloured nuclei of cells (granulations), and especially fungi coloured with aniline or other dyes, become more exquisitely defined. In this way micro-organisms may readily be seen and recognised, even when very sparsely present in a preparation. *The apparatus is indispensable for bacteriological research.**

In addition to a suitable stand and a condenser, good objectives are needed.

First a weak objective, magnifying about 60 to 80 diameters, for the inspection of plate-cultivations; and in addition it is very advantageous to have a good powerful dry objective. There are many objects, as fresh blood, fresh milk, or fresh pus, for the examination of which immersion lenses are not suitable. For such purposes Zeiss's lenses F or D, or Reichert's 8A, may be recommended. With these, especially if a condenser be also used, many bacteriological preparations, as those of tubercle-bacillus from the sputum, may be adequately investigated. For very delicate preparations, and in particular where minute details of structure are to be made out, an immersion-system is needed. The water-immersion systems, formerly much in use, have been superseded of late by the oil-immersion (homogeneous immersion) lenses made by Stephenson and Abbe and Zeiss, which are to be preferred on account of the better definition and clearness of the image which they produce. Instead of water, there is interposed between the front lens of the objective and the object (cover-slip) a fluid having the same refractive index as glass. For this purpose a mixture of fennel and castor-oils may be used. Reichert employed with his lenses a mixture of vaseline and olive-oil, which has the advantage of being odourless, and of penetrating less readily within the lenses. At present concentrated cedar-oil is most in use. Such a system has the further advantage that it does not require correction-collars to be used, as do dry lenses, and that powerful eye-pieces may be employed with it. It is very

* The best results are obtained with the instruments supplied by *Hartnack* (Potsdam), *Seibert* and *Krafft* (Wetzlar), *Leitz* (Wetzlar), and especially by *Zeiss* (of Jena). *C. Reichert* of Vienna supplies with his little microscopes IV. and V. a condenser which is very suitable for clinical purposes.

advantageous also to place a drop of oil on the under surface of the slide which carries the object, between this and the collecting lens of the condenser.

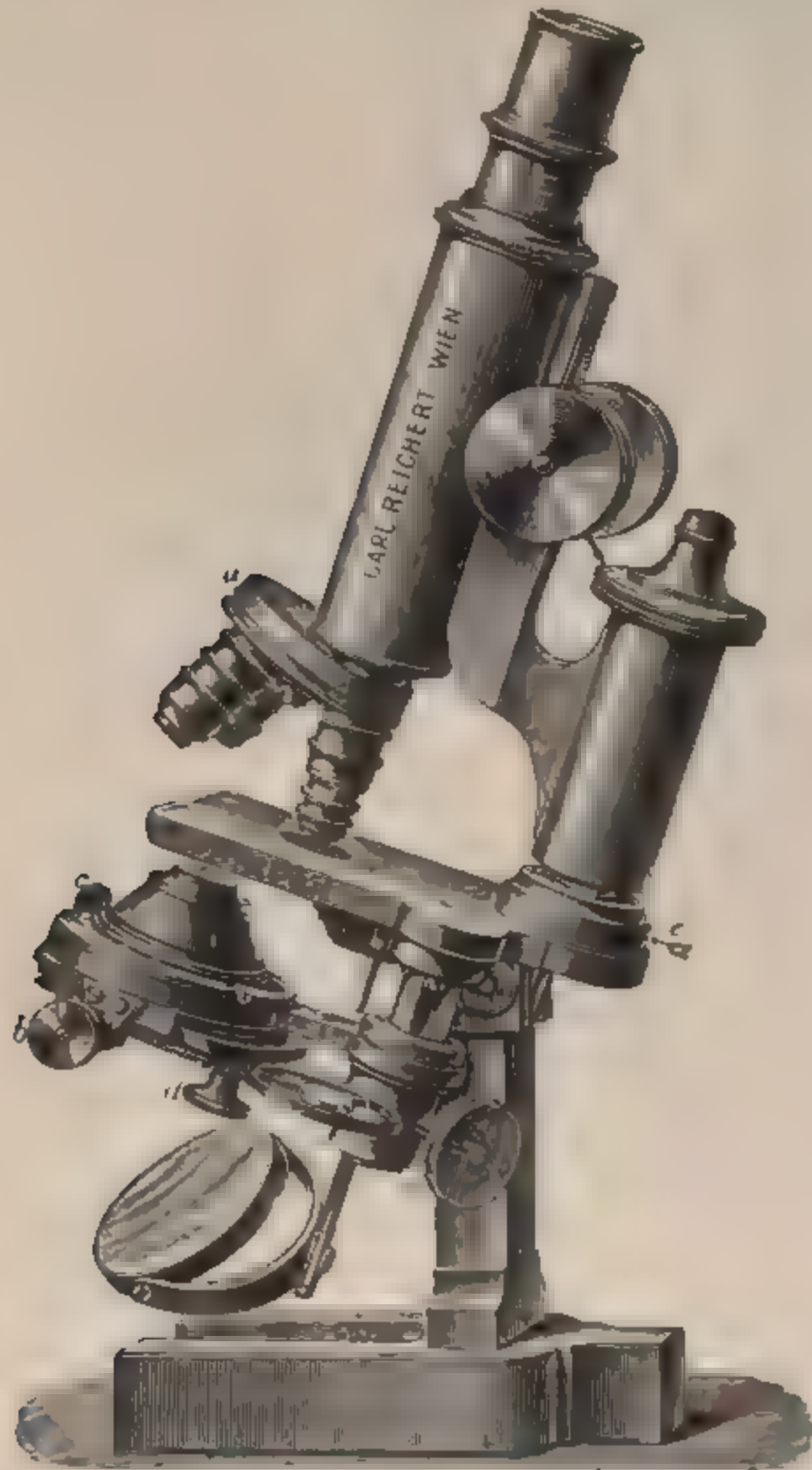


FIG. 155.—Reichert's Stand No. II b., with draw tube track and panier fine adjustment
and Abbe's condenser.

Of less moment is the choice of an eye-piece. In general, for every form of investigation, except bacteriological research, eye pieces of low magnifying power should be taken. For the rest, the eye-pieces II.

and V. as supplied by the firms of Reichert and Zeiss will serve for all cases. The periscopic eye-pieces of Seibert and Krafft are very excellent.*

An objective of crown- and flint-glass was first adopted by Dr. Schott of Jena. It is an admirable contrivance, and has become indispensable for many purposes, as in photographing micro-organisms, where a well-defined achromatic outline is required. Zeiss has given such the name of apochromatic objectives. The corresponding compensation eye-pieces

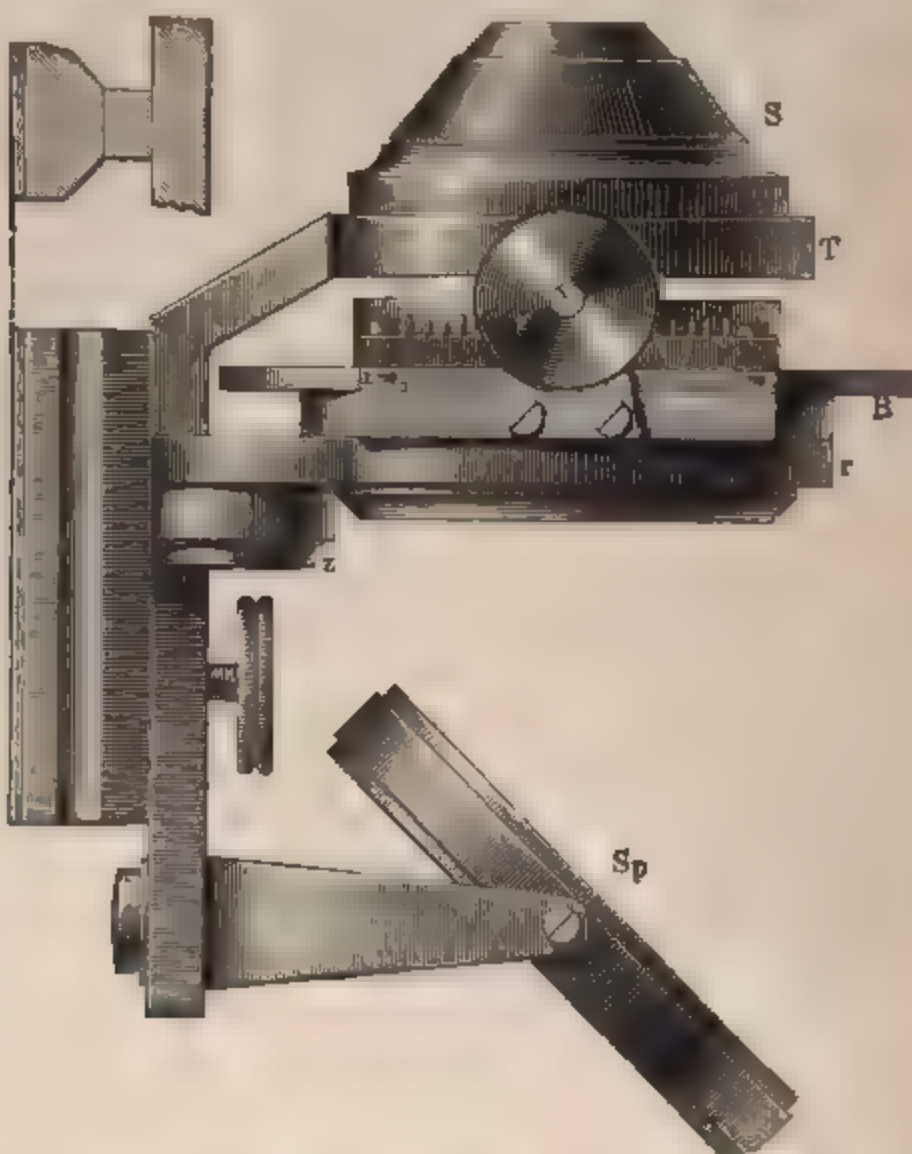


FIG. 156. Abbe's Condenser, as made by Zeiss. Natural size, side view. Sp. Mirror

should be used with them. Their chief advantage lies in the fact that they give a clear and well-defined image with eye pieces of greater strength than could formerly be used. Apochromatic objectives of

* The author has for many years employed an instrument made by Reichert, and he has found it to serve well in every kind of microscopical work, histological and bacteriological. Its parts are as follows.—Eye-pieces II. and IV., objectives 4. 3A, and oil-immersion $\frac{1}{3}$; a small stand with condenser (Abbe's) and cylinder diaphragm. Its price was 207 florins without the oil-immersion, which cost 107 florins. Very good and inexpensive systems of lenses are also made by Plossl of Vienna.

excellent construction are supplied by Reichert. The images of the most delicate objects obtained with Reichert's homogeneous immersion objective of 2 mm. focal length, and even with the working eye-piece 12, are clear and distinct in their smallest details. There is one disadvantage attending the use of these lenses. They need a finer adjustment than the usual mechanism affords, and the image becomes indistinct with the slightest movement of the instrument. It has then

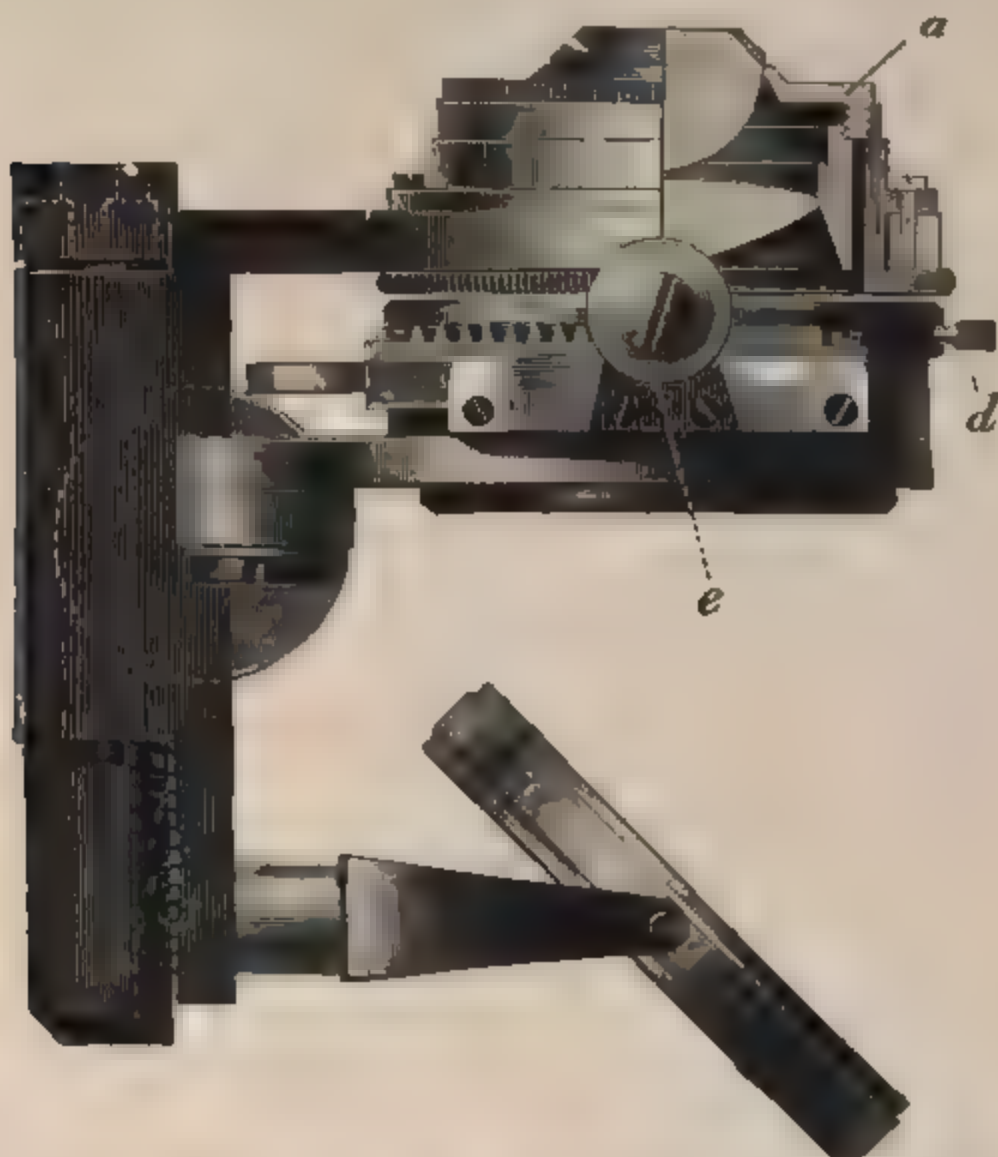


FIG. 157.—Abbe's Condenser *a* above showing the collecting system of lenses.
d Iris diaphragm.

to be focused anew. This defect is apt to produce misleading appearances, and it must be admitted that the otherwise excellent apochromatic objectives supplied by Zeiss are so far in need of improvement. Reichert's hemiaPOCHROMATIC lenses are cheap and easy to work with.

[For bacteriological work, it is an advantage to have a microscope with a stand to which an Abbe's condenser can be fixed under the stage. FIG. 155 shows a convenient form made by Reichert of Vienna. Other makers furnish somewhat similar stands. Abbe's condenser is placed

under the stage, and the tube of the microscope is provided with a "revolver" or "nose-piece," to which lenses of different magnifying powers are fixed.

In an Abbe's condenser (fig. 156) the illuminating apparatus is a condenser system of very short focus (the section of the lenses is shown in fig. 157), which collects the light reflected by the mirror (*Sp*)—the plane side of the mirror being used—into a cone of rays of very large aperture, and projects it on the object. For ordinary work the cone of light is reduced by means of diaphragms, the most convenient form being that known as the "iris diaphragm," which can be adjusted to any size (fig. 157, *d*). Oblique illumination can be obtained by placing the diaphragm eccentrically, which is done by means of the rack-and-pinion movement (*e*).

When very exact definition is required, apochromatic lenses are employed. They are expensive, but the objectives are so constructed as to secure the union of three different colours of the spectrum in one point of the axis. The images projected by them are nearly equally sharp with all the colours of the spectrum. As there is very great concentration of light by these objectives, they permit of the use of very powerful eye-pieces, thus giving high magnifying power with relatively long focal length. A series of *compensating* oculars are used with these lenses. The eye-pieces of extremely low power are called "searchers," while the ordinary or working eye-pieces, beginning with a magnifying power of 4, are classified as 4, 8, 12, 18, and 27. With these eye-pieces great magnifying power is obtained, partly by the lens, but also by the ocular used.]

II. THE DETECTION OF MICRO-ORGANISMS.

In many cases the object to be examined may be placed under the microscope without preparation of any sort. The characteristic micro-organisms will then become visible. This is so with the spirillum of relapsing fever, anthrax-bacilli in the blood, &c. For the most part, however, special processes are necessary for the detection of micro-organisms. The details of some of the processes for the preparation of specimens have been gone into in the chapters on the *Blood*, *Sputum*, &c., which the reader will consult for information concerning them.

It will not be out of place if we give here a brief summary of the methods in use, and point out the particular purposes to which each is applicable. The principles upon which all of them are based were worked out by *Koch*, *Ehrlich*, and *Weigert*; and almost every day some new process, or a modification of the old familiar methods, is made

known. It would far exceed the limits of this book to attempt an account of all the methods which have been suggested, and the various modes of applying them. We shall confine ourselves to what has appeared in the more important and systematic essays dealing with the subject. Those of *Günther*³ and *Unna*,⁴ especially the latter, comprise an accurate and exhaustive account of the methods for staining fungi. Particularly good results in the staining of micro-organisms in sections, which, however, lies beyond our province, have been obtained with *Kühne's*⁵ processes. They have been tested by *Rille*, a pupil of the author, and have been found most valuable for dry cover-glass preparations as well. The methylene-blue method⁶ and *Kühne's*⁷ modification of Gram's method (staining with alcoholic solution of Victoria-blue) are very serviceable in the examination both of sections and of the secretions.

In the examination of the blood and secretions for pathogenic micro-organisms it is in general best to proceed with the use of basic aniline dyes as described in Chapter I. Should it happen that no result is obtained in this way, greater certainty may be secured with *Löffler's* method (see Chapter I.), which is especially suitable for detecting the bacilli of typhoid fever and glanders, and, finally, with Gram's method (see Chapter I.). All the fungi hitherto discovered stain with this, *except the bacilli of typhoid, influenza, and cholera, and gonococci*. The bacillus of hen-cholera is also unaffected by it.

Günther's method (see Chapter I.) is very serviceable for staining the spirillum of relapsing fever. The investigation of the blood and secretions for tubercle-bacillus should be carried out precisely in the manner laid down by *Koch* and *Ehrlich* (see Chapter IV.). Staining with basic aniline dyes serves well for the detection of fungi in the buccal cavity, the nasal secretion, and the gastric contents; but for examining the buccal secretion Gram's or *Günther's* method may also be adopted with advantage, since they render visible the very delicate Spirochæte buccalis (p. 97) and capsulococci. The bacillus of influenza stains best with a weak *Ziehl-Neelsen* (carbol fuchsin) solution. The apparatus devised by *Hofmeister*⁸ is useful for the making of preparations on a large scale.

In searching for the fungi of the alimentary canal, pathogenic and non-pathogenic, all the processes hitherto mentioned should be employed where a thorough investigation is aimed at, and the observer should not forget to add a little iodo-potassic-iodide solution to a drop of the fluid under examination (see p. 201).

In examining the urine, the best results are secured with Gram's or Friedländer's method (p. 131). By their aid the author has detected in various specimens of urine from persons both in health and disease an unexpected profusion of different forms of fission-fungi.

The micro-organisms which occur in pus stain most readily with Gram's method, or its modification already referred to (Victoria-blue). Löffler's and Friedländer's methods are applicable to the same purpose.

To stain the spores of micro-organisms a preparation is made in the manner indicated at p. 46, but exposed for a longer time to the heat. It may be passed through the flame about ten times (*Hueppe*).⁹ When this is done the bacilli lose their staining properties, whilst the spherical objects, if they consist of spores, take up colouring-matter. It is still better to employ the process of double-staining. The preparation is first stained in a hot Ziehl-Neelsen fuchsin solution, then decolorised with nitric acid, and again stained with methylene-blue. The spores then appear red and the bacilli blue.¹⁰

Special processes have been devised for the purpose of defining the flagella which some bacteria present. These have been worked out by *Löffler*, *Künstler*, *Neuhäus*, and *Trenkman*.¹¹ As a mordant *Löffler* employs a fluid composed as follows: solution of tannin (tannic acid 20 parts, water 80 parts) 10 cc.; a cold saturated solution of ferric sulphate, 5 cc.; watery or alcoholic solution of fuchsin, methyl-violet or wool-black, 1 cc. The staining-fluid is neutral saturated anilin-water-fuchsin solution. The method of proceeding will be described here. The cover-glasses are heated with concentrated sulphuric acid, washed with water and then with a mixture of alcohol and ammonia in equal parts, and afterwards polished with a cloth which should be free from grease. Then they are brought in contact with a platinum needle carrying a particle of the pure cultivation, and the latter is spread out finely and divided upon their surface, after which they are allowed to dry in the air. They are next grasped between the fore-finger and thumb of the observer and passed through the flame of a lamp. A drop of the mordant is now supplied to the preparation, the cover-glass is again warmed, and after a minute the preparation is rinsed first with water and then with alcohol. A drop of the staining fluid is next added, and the preparation is heated and washed with water as before.

III. CULTIVATION OF MICRO-ORGANISMS.

A. Methods of Sterilisation.—When the presence of micro-organisms has been ascertained with certainty by the processes described above, the next task is to secure their development outside the system, in other words, to cultivate them. To this end the first requisite is the means of sterilisation. *The essential condition that must be secured in all such cultivation researches is the absolute freedom of the instruments and vessels employed from fungi and germs capable of development.*

In the case of metallic instruments, the necessary cleanliness may be

attained best and most readily by raising them to a red heat in the flame of a Bunsen's burner. Glass vessels, such as test-tubes, flasks, &c., should be purified as far as possible by washing first with distilled water, then with corrosive sublimate solution (1 : 1000), and rinsing out with alcohol and æther, after which they may be sterilised in a dry heat. This can be done best by a sterilising apparatus for temperatures over 200° C. ; but if such an apparatus cannot be had, it may also be done by heating the vessels cautiously over the flame of a Bunsen's burner. In the latter case they should first be carefully dried, in order to prevent the glass from cracking, and the mouth of each should be closed with a compact plug of sterilised cotton-wool before heat is applied.

It is very important to heat the sterilised vessel again immediately before using it, having previously ascertained that the plug is at once sufficiently compact and easily extracted.

The plug may be made of fine glass fibre, or, still better, of asbestos, instead of cotton-wool.

Test-tubes which are to be held in readiness for use are purified in the manner described, fitted with a plug, and then placed in wire baskets, and sterilised in a dry heat.

The sterilisation of the nutrient fluids presently to be mentioned may be effected by boiling them in glass flasks, furnished with a plug of cotton-wool, in those cases where the constitution of the fluid is not altered by heat. To sterilise nutrient gelatine and agar-agar solution (see p. 442), these substances are repeatedly boiled in the vapour sterilisation apparatus. Too frequent, and especially too continued, boiling should be avoided in the case of these two substances, lest they should remain fluid after cooling.

When potatoes are employed as the nutrient substance, they are first freed from sand with a brush, placed for an hour in a 5 per cent. corrosive sublimate solution, finally sterilised (boiled) by steam at boiling-point, and cut into strips with a sterilised knife. Where the vapour sterilisation apparatus devised by *Koch* cannot be had, a Papin's digester with a perforated tray will serve. It is more difficult to sterilise those nutrient substances that will not bear a heat of 100° C. without their parts coagulating and so becoming opaque. In their case, *Koch* recommends that they should be sterilised by intermittent heating. This plan is especially useful for the purpose of freeing blood-serum from fungi and germs.

To prepare sterilised blood-serum, *Koch* proceeds as follows:—The part of the animal's skin through which the blood is to be taken is shaved, and thoroughly cleansed by washing it with solution of corrosive

sublimate, alcohol, and æther. The underlying blood-vessel is freely exposed, and opened with sterilised instruments. The blood is then made to flow directly from the artery in a sterilised glass cylinder, which is filled to the brim and placed in a refrigerator or upon ice for twenty-four to forty-eight hours, so as to allow the corpuscles to settle. The clear amber-coloured serum that has separated after twenty-four hours is drawn off in sterilised pipettes and placed in test-tubes previously sterilised in the manner already indicated. These are heated to 58° C. for two to six hours, and finally the serum is made to coagulate by heating it to 65°–68° C.

With a view to obtaining the largest possible inoculation surface, it is desirable to procure the coagulation of the fluid in the test-tube with its surface inclined very obliquely to the latter. This end may be attained by using a tin vessel with double walls between which water may be retained, covered on top with a plate of glass, and having at its anterior end two movable feet which can be fixed by screws. Or instead of this the test-tube may be fixed by a clamp at an oblique angle in a pot filled with water. For many purposes, and especially for the cultivation of the pathogenic fungi occurring in man, human blood-serum should be employed. To procure human serum the author has adopted the following :—The skin is first thoroughly cleansed in the manner already described, and a puncture is made in it with a cupping-blade previously sterilised by exposure to a heat of 200° C. The blood is taken in a cupping-glass similarly sterilised, and poured into small test-tubes also sterilised. The remainder of the process is carried out as before. Human serum has proved, in the author's experience, to possess notable advantages over that of animals. It remains clearer after coagulation, and is firmer. Where human blood-serum cannot be had, transudation fluid or serous exudation may be taken instead. In that case it is prepared in the same way as the former. A very useful modification of the process for the preparation of blood-serum and of blood-serum plates has been devised by *Unna*.¹² To the blood-serum of the calf, peroxide of hydrogen is added drop by drop, until the mixture, which was at first a brownish-yellow, becomes clear. For this purpose, a quantity equal to about half the volume of the serum will be required. The mixture is neutralised with a 2 per cent. solution of sodium carbonate and passed through a wet filter packed to one-fourth of its depth with well-calciined diatomaceous earth. The fluid which first passes through is usually turbid, and must be refiltered ; and, finally, the clear filtrate is sterilised in the usual manner. For making plate cultivations *Unna* recommends the admixture of 10 per cent. gelatine or 6 per cent. agar-agar.

B. Nutrient Substances.—By the methods described at p. 436 we

are enabled to detect micro-organisms; and the measures that must be taken to render the fluids and nutrient substances as well as the instruments employed in research free from fungi have also been indicated.

It does not suffice to place a fungus or the germs of fungi at random in solid or fluid nutrient substance duly sterilised in order to ensure their successful cultivation; but there is for each of the pathogenic and non-pathogenic fungi an appropriate soil, which differs widely in its chemical constitution in different cases. The researches of *Pasteur*¹³ in connection with yeasts, of *Nägeli*¹⁴ and *Buchner*¹⁵ with bacteria and moulds, of *A. Schultz*¹⁶ with moulds, of the author¹⁷ with the *Micrococcus uræa*, and of *Hueppe*¹⁸ with the lactic acid bacillus, have shown that, in addition to a supply of nitrogen and carbon, each fungus requires certain inorganic salts for its growth. And further, there is in each case some particular temperature at which the fungus thrives best (optimum temperature).

It is only when all these conditions have been secured that cultivation can be made successfully.

To draw unequivocal conclusions from researches of this sort, it is above all necessary to apply Koch's process, presently to be described, for obtaining pure cultivations of the fungus which is being investigated, and to implant these upon solid or fluid nutrient substances.

1. Nutrient Fluids.—The use of fluids as nutrient substances entails uncertainty in the result, since the microscope cannot be employed to examine them. Still it is not difficult to obtain a pure cultivation in a fluid if it be sterilised, and if a cultivation known to be pure be planted in it.

The process is then the same as for the preparation of pure cultivation by Koch's method, which will be described presently.

The composition of the nutrient fluid must vary with the nature of the fungus to be cultivated.

Thus yeasts grow best in a somewhat acid saccharine fluid. Moulds require a medium holding free acids in considerable quantity. Feebly alkaline fluids are the best for certain non-pathogenic bacteria. Fluids of definite constitution for the cultivation of bacteria—as those of *Pasteur*, *Cohn*, and the author—agree in that they all contain nitrogenous and carbonaceous matter and inorganic salts.

Although it is true that by cultivation in nutrient fluids much useful information has been obtained concerning the life-history of some of the fission-fungi, the method is not employed for the investigation of pathogenic micro-organisms, partly because, as has been said already, there always remains some doubt as to whether the cultivations obtained with them are in reality pure cultivations, and partly because it would seem that pathogenic fungi thrive badly in fluids. The author has en-

endeavoured, without success, to obtain pure cultivations of pneumonia-cocci, Streptococcus pyogenes aureus, and other pathogenic micro-organisms, on sterilised nutrient fluids of the most varied constitution.

Control experiments have proved that non-pathogenic fungi developed readily in such nutrient fluids, while the same fluids, maintained under like conditions, when infected with pathogenic fungi remained sterile.¹⁹

2. Solid Nutrient Substances.—As in the case of nutrient fluids, the chemical constitution of solid nutrient substances will vary greatly according to the biological character of the fungus under cultivation.

1. Blood-Serum.—The blood-serum of animals is required for certain pathogenic fungi, e.g., tubercle-bacillus; gonococci are cultivated in human serum. The method of preparation has been sufficiently described.

2. R. Koch's Peptone-gelatine Bouillon.—This is prepared in the following manner:—500 grms. of good meat, free from fat and freshly minced, is rubbed up with 1000 grms. of distilled water and placed for twenty-four hours in a refrigerator. It is then strained through linen, the resulting fluid made up to 1000 cc., and 10 grms. of dry peptone, 5 grms. of common salt, and 100 grms. of white edible gelatine are added. The fluid is heated in a flask until the gelatine is dissolved. It is then accurately neutralised with sodium carbonate, and boiled for half-an-hour to an hour, when a specimen is tested for its reaction; after this the fluid is filtered through a hot-water funnel and poured into test-tubes sterilised in the manner directed (see p. 439), and sterilised for ten minutes a day for three days.

The test-tubes may now be kept for weeks or months at the ordinary temperature before they are used; but when this is done, an india-rubber cap should be placed over each above the plug of cotton-wool, to prevent evaporation from the gelatine. Keeping the nutrient gelatine thus for a long time in the test-tubes has this advantage, that when through an accident in its preparation germs have obtained admission to the fluid, the resulting cultivations are made evident by clouding in the gelatine, which will not then be employed for cultivation research.

Koch's nutrient gelatine, which may be modified at will by the addition of organic or inorganic substances, will serve for the cultivation of all pathogenic and non-pathogenic fungi which grow at the ordinary temperature of a room. It cannot be used at higher temperatures (over 25°-30° C.), which cause it to melt, and for micro-organisms which liquefy gelatine rapidly.

3. Agar-Agar.—In many investigations, especially for fungi which require a temperature not lower than blood-heat to develop well, or which liquefy gelatine rapidly, agar-agar may be used with advantage

instead of gelatine as a nutrient substance. It is prepared in the same way as nutrient gelatine fluid, only that instead of gelatine 1.5–2 per cent. of finely divided agar-agar is added to the fluid. Unfortunately it is difficult to prepare a perfectly pure and clear agar-agar solution in this way, and that substance, even when added in small particles, filters badly through the hot-water funnel. *Schottelius*²⁰ and *Richter*,²¹ however, have modified the process in such a manner that a clear nutrient agar-agar may be easily obtained.

4. Potato.—Enough has been said about the sterilisation of potatoes employed for nutrient material (p. 439). Their use is of great service in studying pathogenic fungi, many of which develop on potato in a highly characteristic manner (see pp. 206, 411).

Starch to which suitable nutrient salts have been added forms a solid nutrient substance, which is very useful and easily sterilised. Together with gluten and bread it is especially good for the cultivation of moulds. For the latter also boiled blood clots may be employed with advantage.²² Both starch and blood may be easily and effectually sterilised by a current of steam.

Recent investigators have in many instances somewhat modified the constitution of, and the method of preparing, nutrient substances. Thus the addition of glycerine to peptone gelatine or agar-agar has been tried successfully. All the modifications suggested cannot be noticed here.²³ The use of stained nutrient solids and fluids, however, demands notice.

According to *Birch-Hirschfeld*,²⁴ living anthrax-bacilli can be procured stained by inoculating from a pure cultivation of the fungus upon 15 per cent. nutrient gelatine which contains in 6 cc. 1 cc. of a watery solution of fuchsin or methylene-blue. The cultivation should remain for twenty-eight hours at 35°–40° C.²⁵

In the cultivation of the bacillus of typhoid fever also, stained (benzo-purpurin) nutrient substances are useful.

The application of this principle, as by the addition of neutral tincture of litmus,²⁶ or other substances, to show the formation of free acid or acid salts, has thrown much light upon the nature of pathogenic and non-pathogenic fungi. The addition of a little watery solution of Congo-red or benzo-purpurin to nutrient substances has proved, in the author's hands, a valuable expedient in the study of the biological character of micro-organisms. If the material examined contains acid-forming fungi, the cultivations upon stained nutrient soil take a colour ranging from a pale to a blackish blue, and their presence is discernible with the naked eye. The method is particularly appropriate for the study of micro-organisms from the intestine under normal and pathological conditions.

C. Preparation of Koch's Pure Cultivations.—Although *Klebs*²⁷ and *Brefeld*²⁸ had already suggested, and themselves adopted, the use of solid nutrient substances for the investigation of fungi, it was *Koch* who grasped the significance of these methods, and, by submitting the cultivations in fluid and solid media to the control of the microscope, laid the foundation of modern bacteriology.

To this observer we owe not only the knowledge of many fundamental facts in bacteriology, such as result from the discovery of the bacillus of tubercle and of cholera, but nearly all the cultivation and staining methods in use are derived from his researches and those of his pupils.

The methods now to be described have for their object as far as possible to separate the individual germs from a mass of fungi, and to promote their development apart from one another in solidifying fluids. This may be done by Koch's processes for obtaining plate-cultivations on glass slides and test-tube cultivations. It is usually expedient to make plate and test-tube (deep inoculation) cultivations together.

1. Plate-Cultivations.—A test-tube, in which from 5–8 cc. of solidified nutrient gelatine has been placed as directed above, is put into warm water, and allowed to remain there until the gelatine is quite fluid. Care is now taken that the plug in the mouth of the test-tube is freely movable by rotating it a little if necessary. The test-tube is held obliquely between the thumb and index-finger of the left hand, the plug with the upper end between the second and third fingers (Koch's "grasp"). A little of the fungoid material is taken with a freshly-sterilised platinum needle, and placed on the gelatine, so as first to touch its edge and then to mix with the fluid. In doing this, draughts of air must be excluded from the room. In the same way one or more drops of this first dilution are placed in nutrient gelatine in another test-tube (second dilution), and if a provisional inspection has shown that the fluid under examination is very rich in fungi, the process is repeated again (third dilution). It may then be assumed that the germs in the nutrient gelatine are actually isolated. The infected gelatine is next poured out on to glass plates about 14 cm. in length and 12 cm. broad, and caused to solidify quickly. This may be done in a few minutes by the application of cold. The glass plates are prepared in the following manner:—They are first thoroughly cleansed with water, solution of corrosive sublimate, and alcohol, and placed immediately before use in iron cages in the steriliser, where they are heated for a long time at 100°–150° C. They are taken out when cool, and laid upon a large sheet of plate-glass made cold with ice. This should have a polished, even surface, and must also be accurately level when used, an end best attained by mounting it on a tripod stand with a spirit-level

visible beneath the glass. Such an accessory, however, is not essential, and it may be dispensed with if a little care be taken.

Recently the use of the glass plate cooled on ice has been supplanted in many laboratories by that of a polished plate of iron about 20 cm. in diameter and 8 cm. thick, which is carefully sterilised beforehand. This is placed upon the tripod and adjusted with the spirit-level. An iron plate was first employed in Lichtheim's laboratory at Berne, and it is very suitable to the purpose in hand, because with it nutrient fluids solidify very quickly. In warm weather, as in the summer, the iron plate must be previously cooled on ice, but ordinarily this precaution is unnecessary. The author has employed such a plate exclusively in his bacteriological work during the last four years, and found it most serviceable.

The nutrient gelatine is poured upon the plates in the following manner:—The small glass plates upon which nutrient gelatine is to be spread are laid upon the glass plate made cold with ice or on the iron plate, and that edge of the test-tube over which the gelatine is to flow is heated. When the edge has cooled, the gelatine is poured here and there over the cold surface of the plates, and spread out upon them as evenly as possible by means of the sterilised edge of the test-tube, care being taken that the edges of the plates are left free. The plates are then covered with a bell-glass. When solidification is complete, they are placed on sterilised wet blotting-paper in a glass dish of about 30 cm. diameter, which has been previously well purified with corrosive sublimate, and another bell-glass is put over them.

In order to obtain the sterilised and wet blotting-paper, a jet of super-heated steam is directed upon the filter-paper in the bottom of the glass dish for from eight to ten minutes. In this way the bell-jar as well as the filter-paper are sterilised, and the latter is saturated with sterilised vapour.

In such a vessel six plates, or even more, can be put, with a glass partition between each pair.

Instead of plates, *E. Esmarch*²⁹ has recently employed test-tubes, which will serve for many of the purposes of plate-cultivations. With these the process is as follows. Some of the fungoid material is placed in the fluid nutrient gelatine in the test-tube in the manner already described, and mixed with it as intimately as possible. The test-tube, covered with an india-rubber cap above the plug of cotton-wool, is then held as vertically as possible, and with the sterilised plug and cap directed somewhat upwards, under a stream of cold water, and in this position is made to rotate upon its long axis. After a little while the nutrient gelatine has solidified in the cylindrical form of the test-tube. This process has considerable advantages. Cultivations so made can be inspected not only with weak (Reichert IV.), but also with powerful

objectives ; they are less likely to be contaminated ; individual cultivations may be removed with a little care even under the microscope, and less annoyance is apt to arise from the unpleasant odour which the cultivations often give rise to than is the case when plates are used. To these the name of cylinder-cultivations may be appropriately given. On a plate or in a test-tube so prepared, after a longer or a shorter interval, there appear little punctiform colonies, which are already sufficiently distinguishable from one another, and at the same time the gelatine is often partly liquefied and gives off a disagreeable smell. If now a minute particle be taken from each of the colonies by means of a sterilised platinum needle and submitted to the same process over again, pure cultivations will be had of all the fungi which develop in nutrient gelatine.

At the same time, by placing the entire plate under the microscope and examining it, it will be possible to study the mode of growth of the fungi, and also to determine whether in any given case the cultivations are pure or contaminated by admixture with other fungi. Certain distinctions in the shape and colour of the cultivations can also be recognised with the naked eye, and by removing a particle on a platinum needle from an individual developing cultivation under the microscope and transferring it directly to a test-tube (cultivation by deep inoculation), a definite fungus may be cultivated in a little time and with little trouble.

Cultivations with nutrient agar-agar are made in the same way as with nutrient gelatine. The former substance should be used in all cases of fungi which contain spores and where the growing fungi cause nutrient gelatine to liquefy quickly ; as, for instance, in making cultivations from the fæces, and in cultivating micro-organisms which require a high temperature (37° and upwards) for their growth.

The cultivations are placed in an incubator such as Koch and others (d'Arsonval) have devised. The construction of these is the same in all cases. The incubator has double walls enclosing a compartment for water, with arrangements (thermostats) by which the internal temperature may be maintained constant to within 0.2° C. Experience, and especially Koch's researches, have shown that a number of pathogenic fungi, as, for instance, the bacillus of tubercle, will thrive only at a definite and continuous temperature. To aid in securing this, various thermostats have been constructed recently. Of these, the author uses the thermo-regulator of *L. Meyer*.³⁰ The principle of the instrument is to regulate the supply of gas for heating the incubator by conducting it through an æthereal atmosphere confined by a mercurial valve, so that more or less gas is conducted to the jet according to the temperature required in the incubator.

This instrument answers well. Such a thermostat has been in use for several months in Professor Nothnagel's clinic. Notwithstanding the great variation in gas-pressure, the temperature recorded never varied by more than 0.2° C.

2. Cultivation by Deep Inoculation.—A particle of the fungoid mass is taken on a sterilised platinum needle and implanted on nutrient gelatine or agar-agar in a test-tube by removing the sterilised plug with the mouth of the tube downwards and plunging the inoculating needle into the nutrient substance. After the lapse of a few days the fungus develops in a very characteristic manner in the gelatine. This proceeding is adopted only when a pure cultivation has been obtained with plates, and it is then of service in the further study of the micro-organism. *R. Fischl*³¹ has removed the cylinder of gelatine from the test-tube with a cork-borer, and *Neisser*³² by the action of heat; and by subsequently hardening it in alcohol or a 1 per cent. solution of bichromate of potash, these observers were able to observe the development of the fungus in microscopical sections.

3. Cultivations on Glass Slides.—A sterilised platinum needle is infected with a particle from the fungoid fluid in the manner indicated on p. 444, and drawn across the surface of nutrient gelatine spread upon a slide in such a manner that the fungi lodge in the furrow. After a few days colonies of fungi develop in the line of inoculation.

4. Cultivation in Hanging Drops.—*Koch* was the first to adopt this method of cultivation. With it the growth of the micro-organisms can be observed directly under the microscope. It is carried out thus:—A glass slide with a hollow surface is taken, and the edge of the concavity is smeared with a little vaseline, or a mixture of five parts of vaseline and one of paraffin may be used instead, as *Birch-Hirschfeld* recommends. A drop of sterilised broth is then placed on a clean cover-glass infected from the bacterial fluid, and covered with the slide in such a way that the drop is suspended in the middle of the cell. The sterilised broth is prepared in the same manner as nutrient gelatine, except that the addition of gelatine is omitted. For microscopical examination an oil-immersion lens with an *Abbe's* condenser and narrow diaphragm should be used. And where fungi are studied in this way, the edges of the drop should be observed with special care, since it is there that the morphological characteristics of the micro-organisms are best marked.

5. Cultivation by Exclusion of Air.—A number of micro-organisms develop only in the absence of air (oxygen), and the processes for their cultivation have been worked out by *Koch*, *Hesse*, *Buchner*, *Gruber*, and others.

Koch seals up the test-tube cultivation with plates of mica, *Hesse*

with oil. *Gruber* exhausts the air with an air-pump and fuses the vessel. *Buchner*³³ employs a solution of pyrogallol and caustic potash to absorb oxygen. When this is to be done, the test-tube cultivation, suitably prepared, is placed within a second larger test-tube, which contains the solution, and which is then closed at the top with an air-tight caoutchouc cap. For cultivations in hanging drops, *Buchner's* method is recommended by *Nikiforoff*.³⁴ The exclusion of air from plate-cultivations is best effected by *Blücher's*³⁵ contrivance.

IV. THE TRANSMISSION OF PURE CULTIVATIONS TO ANIMALS.

This constitutes a very important part of bacteriological study. It may be conducted in many ways.

(a.) The animal is placed in a closed chamber, and the atmosphere is saturated with sterilised water containing the bacteria by means of a spray producer. Experiments of this kind are very valuable for the study of infectious diseases and of inhalation remedies.

(b.) The pure cultivation of a definite fungus is given to the animal with its food. In doing this the chief precaution necessary is to see that the food itself is innocuous. *Koch's* plan is to enclose the cultivation in a small starch capsule provided with a lid, and to place this on the back of the animal's tongue. Most of the bacteria that are devoid of spores appear to be destroyed by the action of the free acid in the stomach, and on this account it is advisable to neutralise the acid by the administration of alkalies, such as *Koch* used in his cholera researches, or to perform laparotomy with strict anti-septic precautions, and to introduce the cultivation directly within the duodenum.

(c.) Cutaneous inoculation. The hair having been removed from some part of the body, as the ear, which the animal cannot easily reach with its tongue, a superficial wound is made there, and a portion of the cultivation is lodged within it.

(d.) In mice the inoculation is best made subcutaneously at the root of the tail. Or the purpose may be effected by infecting subcutaneously, or within one of the natural cavities, by means of *Koch's* modification of *Pravaz's* syringe. In this instrument a disc of cork is substituted for the india-rubber, which will not bear the great heat necessary to ensure sterilisation. A part of the cultivation suspended in water is sucked into the syringe, and the fluid injected beneath the animal's skin. A simple glass cannula with a compressible india-rubber bulb will also serve.

V. SCHEME OF A BACTERIOLOGICAL INVESTIGATION.

1. The fluid to be examined is taken from the body with sterilised instruments and under the requisite precautions, and a drop is inspected microscopically, either with a powerful dry system or a homogeneous immersion objective; narrow diaphragms and an Abbe's condenser being also used. Dry preparations are made and stained. In doing this, solutions of the basic aniline dyes are used, or one of the other methods (p. 436), as those of Gram, Friedländer, &c., according to the character of the fungi supposed to be present.

2. Another drop of the fluid is added to fluid nutrient gelatine or agar-agar to obtain plate-cultivations.

The plate-cultivations are examined with the microscope after twelve to twenty-four hours, and it is seen whether they contain fungi which are identical with, or resemble in their mode of growth, others already known or under observation.

Should it appear that no pure cultivations have yet formed, other plates are made from those already prepared, until one is obtained in which a single fungus develops.

3. Drop-cultivations are made, and their growth observed directly. Moreover, these are nourished on various substances, such as potatoes, gluten, &c., and their behaviour to different nutrient fluids and under different conditions of temperature is studied.

4. Pure cultivations so obtained are transferred to different animals, and the morbid symptoms induced are observed. If such symptoms are analogous to those occurring in the human system in presence of the same micro-organism, they may be assumed to be due in both cases to the agency of the latter.

When this is done, however, the resources of bacteriology are not yet exhausted. The biological characteristics of the fungi remain to be investigated as to what sources of nitrogen, what sources of carbon, and what inorganic salts they require. It is only in this way that we can arrive at just conclusions concerning the nature of infectious diseases, and attain to rational methods of anti-bacterial treatment.

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CHAPTER I

THE BLOOD

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CHAPTER IV

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CHAPTER V

GASTRIC JUICE AND VOMIT

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CHAPTER VI

THE FÆCES

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CHAPTER VII

THE URINE

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CHAPTER VIII

EXUDATIONS, ETC.

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CHAPTER IX

SECRECTIONS OF GENITAL ORGANS

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CHAPTER X

BACTERIOLOGICAL RESEARCH

¹ The following list includes the most important works on Bacteriology, with special reference to those which describe the methods of examination and the morphology of Bacteria. We direct attention, as a basis of study, above all, to the already-mentioned works of *R. Koch* and his school.—*Crookshank*, Chap. iv. (54).—*Flügge*, Chap. i. (195).—*Cornil* and *Babes*, *Les Bactéries*. Paris, 1885.—*C. Fränkel*, *Grundriss der Bakterienkunde*. Berlin, 1887.—*A. Johne*, *Ueber die Koch'schen Reinculturen und die Cholerabacillen*. Leipzig, 1885.—*W. Zopf*, *Die Spaltpilze*, 3rd edit. Berlin, 1885.—*C. Friedländer*, *Mikroskopische Technik*, 3rd edit., 1885.—*Siebenmann*, *Die Fadenpilze*. Wiesbaden, 1883.—*A. de Bary*, *Vergleichende Morphologie und Biologie der Pilze*, Leipzig, 1884; and *A. de Bary*, *Vorlesungen über Bakterien*, Leipzig, 1885.—*K. Huber* and *A. Becker*, *Die pathologisch-histologischen und bacteriologischen Untersuchungsmethoden*. Leipzig, 1886.—*H. Mittenzweig*, *Die Bakterien-Aetiologie der Infectionskrankheiten*. Berlin, 1886.—*Duclaux*, *Le Microbe et la Maladie*. Paris, 1886.—*Gottstein*, *Die Verwertung der Bakteriologie in der klinischen Diagnostik*. Berlin, 1887.—*Baumgarten*, *Lehrbuch der pathol. Mykologie*. Brunswick, 1886 to 1888.—*Löffler*, *Vorlesungen über die geschichtliche Entwicklung der Lehre von den Bakterien*. Leipzig, 1887.—*Hueppe*, *Die Methoden der Bakterienforschung*, 5th edit. Wiesbaden, 1891.—*Günther*, *Einführung in das Studium der Bakteriologie*, 2nd edit. Leipzig, 1891.—*Axel Holst*, *Uebersicht über die Bakteriologie für Aerzte und Studierende*, German translation by *Reyher*. Basel, 1891.—*Eisenberg*, *Bakteriologische Diagnostik*. Hamburg and Leipzig, 1891.—*Heim*, *Lehrbuch d. bakteriologischen Untersuchung u. Diagnostik*. Stuttgart, 1894.—*Schenk*, *Grundriss der Bakteriologie*. Vienna and Leipzig, 1894.

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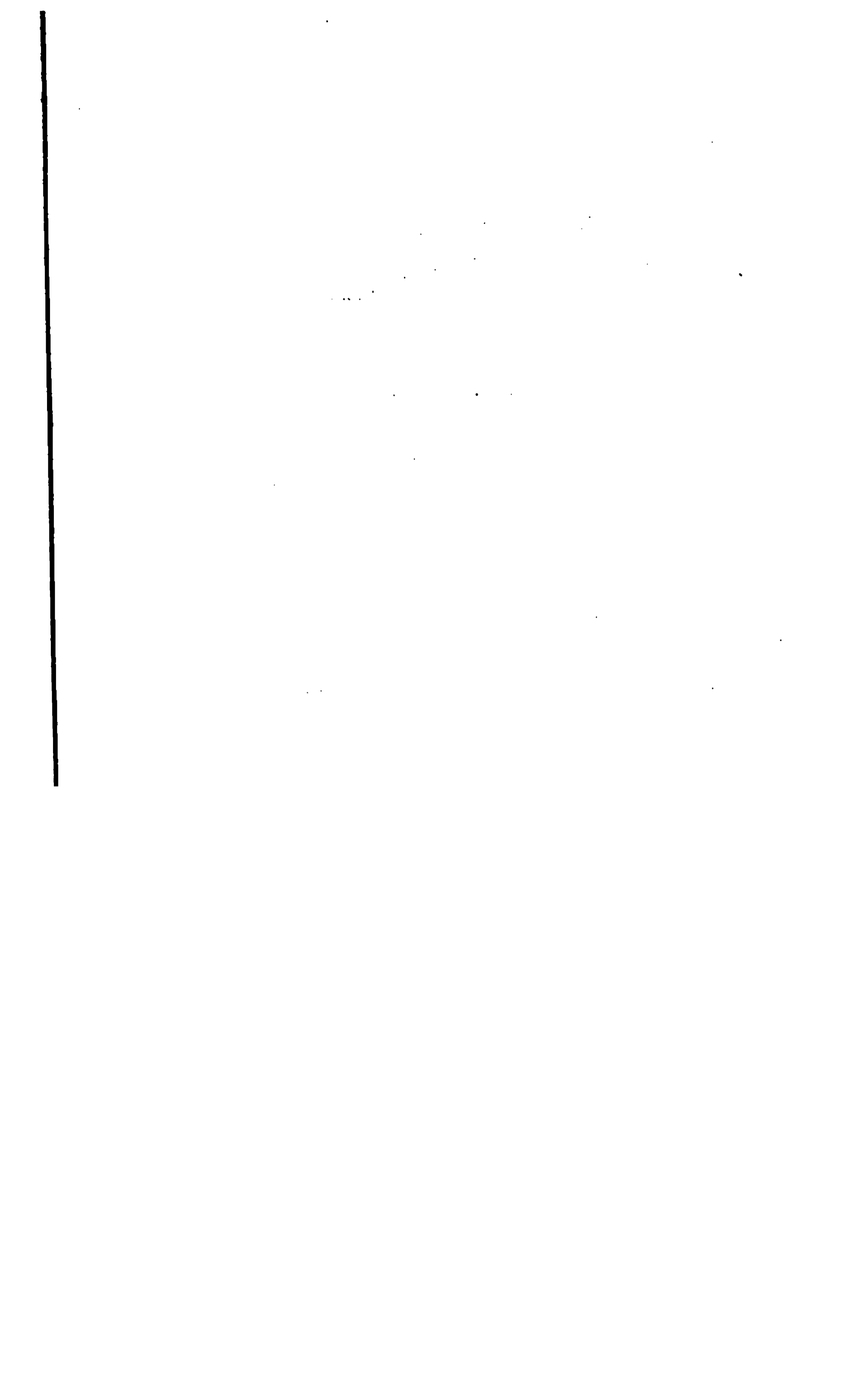
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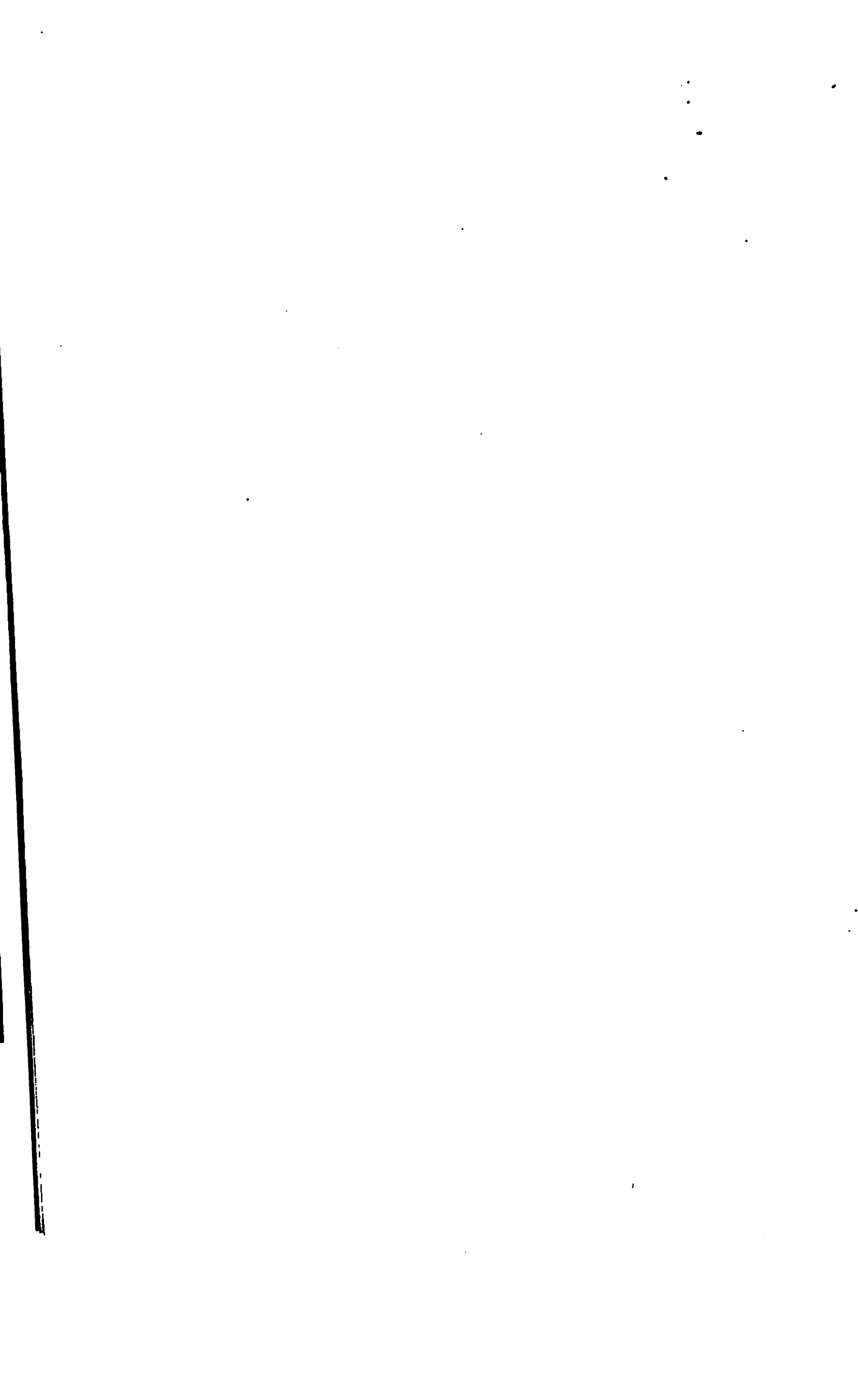
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